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(57) Abstract	54) Title: GREEN FLUORESCENT PROTEINS FOR MEASURING THE PH OF A BIOLOGICAL SAMPLE 57) Abstract							
Disclosed are fluorescent protein sensors for meas	the gre	he pH of a sample, nucleic acids encoding them, and methods of use. en fluorescent protein (GFP) from Aequora victoria. Also disclosed are gion of a cell, such as the mitochondrial matrix or the Golgi lumen.						

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GREEN FLUORESCENT PROTEINS FOR MEASURING THE PH OF A BIOLOGICAL SAMPLE

Statement as to Federally Sponsored Research
This invention was made with Government support under Grant
No. NS27177, awarded by the National Institutes of Health.
The Government has certain rights in this invention.

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Field of the Invention

The invention relates generally to compositions and methods for measuring the pH of a sample and more particularly to fluorescent protein sensors for measuring the pH of a biological sample.

Background of the Invention

The pH within various cellular compartments is regulated to provide for the optimal activity of many cellular processes. In the secretory pathway, posttranslational processing of secretory proteins, the cleavage of prohormones, and the retrieval of escaped

20 luminal endoplasmic reticulum proteins are all pH-dependent. Several techniques have been described for measuring intracellular pH. Commonly used synthetic pH indicators can be localized to the cytosol and nucleus, but not selectively in organelles other than those in the endocytotic pathway.

25 In addition, some cells are resistant to loading with cellpermeant dyes because of physical barriers such as the cell wall in bacteria, yeast, and plants, or the thickness of a tissue preparation such as brain slices.

Several methods have been described for measuring pH 30 in specific regions of the cell. One technique uses microinjection of fluorescent indicators enclosed in

liposomes. Once inside the cell, the liposomes fuse with vesicles in the trans-Golgi, and the pH of the intracellular compartments is determined by observing the fluorescence of the indicator. This procedure can be laborious, and the fluorescence of the indicator can be diminished due to leakage of the fluorescent indicator from the Golgi, or flux of the fluorescent indicator out of the Golgi as part of the secretory traffic in the Golgi pathway. In addition, the fusion of the liposomes and components of the Golgi must take place at 37°C; however, this temperature facilitates leakage and flux of the fluorescent indicator from the Golgi.

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A second method for measuring pH utilizes retrograde transport of fluorescein-labeled verotoxin 1B, which stains the entire Golgi complex en route to the endoplasmic reticulum. This method can be used, however, only in cells bearing the receptor globotriaosyl ceramide on the plasma membrane, and it may be limited by the residence time of the verotoxin in transit through the Golgi.

In a third method, intracellular pH has been measured using the chimeric protein CD25-TGN38, which cycles between the trans-Golgi network and the plasma membrane. At the plasma membrane, the CD25-motif binds extra-cellular anti-CD25 antibodies conjugated with a pH-sensitive

25 fluorophore. Measurement of fluorescence upon return of the bound complex to the Golgi can be used to measure the pH of the organelle.

Summary of the Invention

- 2 -

The invention is based on the discovery that proteins derived from the Aequora victoria green fluorescence protein (GFP) show reversible changes in fluorescence over physiological pH ranges.

Accordingly, in one aspect, the invention provides a method for determining the pH of a sample by contacting the sample with an indicator including a first fluorescent protein moiety whose emission intensity changes as the pH varies between 5 and 10, exciting the indicator, and the determining the intensity at a first wavelength. The emission intensity of the first fluorescent protein moiety indicates the pH of the sample.

In another aspect, the invention provides a method

for determining the pH of a region of a cell by introducing
into the cell a polynucleotide encoding a polypeptide
including a first fluorescent protein moiety whose emission
intensity changes as the pH varies between 5 and 10,
culturing the cell under conditions that permit expression
of the polynucleotide, and determining the intensity at a
first wavelength. The emission intensity of the first
fluorescent protein moiety indicates the pH of the sample.

In a further aspect, the invention provides a functional engineered fluorescent protein whose amino acid sequence is substantially identical to the amino acid sequence of the 238 amino acid Aequora victoria green fluorescence protein shown in FIG. 3 of USSN 08/911,825 (SEQ ID NO:), and whose emission intensity changes as pH varies between 5 and 10.

In another aspect, the invention provides a polynucleotide encoding the functional engineered fluorescent protein.

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The invention also includes a kit useful for the detection of pH in a sample, e.g., a region of a cell. The kit includes a carrier means containing one or more containers comprising a first container containing a polynucleotide encoding a polypeptide including a first

fluorescent protein moiety whose emission intensity changes $\bar{\ }$ as the pH varies between 5 and 10.

Brief Description of the Drawings

FIG. 1 is a schematic diagram depicting fluorescent protein sensors used as indicators of intracellular pH. 5 FIGS. 2A and 2B are graphs showing absorbance as a function of wavelength for the fluorescent protein pH sensor EYFP (SEQ ID NO:__) at various wavelengths (FIG. 2A), and the pH dependency of fluorescence of various GFP fluorescent protein sensors in vitro and in cells (FIG. 2B). The 1.0 fluorescence intensity of purified recombinant GFP mutant protein (solid symbols) as a function of pH was measured in a microplate fluorometer. The fluorescence of the Golgi region of HeLa cells expressing proteins having the 81 Nterminal amino acids of the type II membrane-anchored 15 protein galactosyltransferase (GT:UDP-galactose-β,1,4galactosyltransferase, EC 2.4.1.22) ("GT") fused to EYFP, or EGFP, i.e., GT-EYFP or GT-EGFP (open symbols) was determined during pH titration with the ionophores monensin/nigericin in high KCL solutions. 2.0

FIGS. 3A and 3B are graphs showing ratiometric measurements of pH_0 by cotransfecting HeLa cells with polynucleotides encoding GT-ECFP and GT-EYFP. FIG. 3A is a graph showing single wavelength fluorescence intensities of GT-EYFP and GT-ECFP in the Golgi region of a HeLa cell. FIG. 3B is a graph showing the ratio of GT-EYFP/GT-ECFP fluorescence in the same cell as a function of time.

FIG. 4 is a graph showing the change in mitochondrial pH of HeLa cells expressing YFP H148G.

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FIG. 5 is a graph showing the mitochondrial pH of chick skeletal myotubes expressing YFP H148G in the mitochondrial matrix.

FIG. 6 is a graph showing fluorescence and pH in HeLa cells expressing YFP H148Q targeted to the mitochondrial matrix.

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FIG. 7 is a graph showing a ratiometric measurement of mitochondrial pH following expression of YFP H148G (pH sensitive) and GFP T203I (pH insensitive) in mitochondria of HeLa cells.

FIG. 8 is a graph showing normalized absorbance of WT GFP and YFP in 75 mM phosphate pH 8.0, 140 mM NACl. Solid lines, WT GFP; Dashed lines, YFP.

FIG. 9 is a stereoview of the 2F₀-F_c electron density 15 map of the YFP chromophore and the stacked Tyr203 after refinement. The 2.5 Å resolution map was contoured at +1 standard deviation.

Detailed Description

The invention provides genes encoding fluorescent sensor proteins, or fragments thereof, whose fluorescence is sensitive to changes in pH at a range between 5 and 10. The proteins of the invention are useful for measuring the pH of a sample. The sample can be a biological sample and can include an intracellular region of a cell, such as the lumen of the mitochondria or golgi. The pH of a sample is determined by observing the fluorescence of the fluorescent sensor protein.

The fluorescent protein pH sensor have a broad applicability to cells and organisms that are amenable to gene transfer. Problems associated with the use of other agents used to measure pH, e.g., problems associated with permeabilizing cells to ester-containing agents, leakage of agents, or hydrolysis of agents are avoided. With the

fluorescent protein pH sensors of the invention, no leakage occurs over the course of a typical measurement, even when the measurement is made at 37°C.

Compositions and methods described herein also avoid
the need to express and purify large quantities of soluble
recombinant protein, purify and label it in vitro,
microinject it back into cells. An important advantage of
the fluorescent protein pH sensors of the invention is that
they can be delivered to cells in the form of
polynucleotides encoding the protein sensor fused to a
targeting signal or signals. The targeting signal directs
the expression of the protein sensors to restricted cell
locations. Thus, it is possible to measure the pH of a
precisely defined cellular region or organelle.

15 POLYNUCLEOTIDES AND POLYPEPTIDES

In a first aspect, the invention provides a functional engineered fluorescent protein whose amino acid sequence is substantially identical to the 238 amino acid Aequora victoria green fluorescence protein shown in Fig. 3 of USSN 08/911,825 (SEQ ID NO:). The term "fluorescent protein" refers to any protein capable of emitting light when excited with appropriate electromagnetic radiation, and which has an amino acid sequence that is either natural or engineered and is derived from the amino acid sequence of Aequorea-related fluorescent protein. The term "fluorescent protein pH sensor" refers to a fluorescent protein whose emitted light varies with changes in pH from 5 to 10.

The invention also includes functional polypeptide fragments of a fluorescent protein pH sensor. As used

30 herein, the term "functional polypeptide fragment" refers to a polypeptide which possesses biological function or activity which is identified through a defined functional

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assay and which is associated with a particular biologic, morphologic, or phenotypic alteration in the cell. The term "functional fragments of a functional engineered fluorescent protein" refers to fragments of a functional engineered protein that retain a function of the engineered fluorescent protein, e.g., the ability to fluoresce in a pH -dependent manner over the pH range 5 to 10. Biologically functional fragments can vary in size from a polypeptide fragment as small as an epitope to a large polypeptide.

Minor modifications of the functional engineered fluorescent protein may result in proteins which have substantially equivalent activity as compared to the unmodified counterpart polypeptide as described herein. Such modifications may be deliberate, as by site-directed mutagenesis, or may be spontaneous. All of the polypeptides produced by these modifications are included herein as long as the pH-dependent fluorescence of the engineered protein still exists.

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A functional engineered fluorescent protein includes amino acid sequences substantially the same as the sequence set forth in SEQ ID NO:__, and whose emission intensity changes as pH varies between 5 and 10. In some embodiments the emission intensity of the functional engineered fluorescent protein changes as pH varies between 5 and 8.5.

By "substantially identical" is meant a protein or polypeptide that retains the activity of a functional engineered protein, or nucleic acid encoding the same, and which exhibits at least 80%, preferably 85%, more preferably 90%, and most preferably 95% homology to a reference amino acid or nucleic acid sequence. For polypeptides, the length of comparison sequences will generally be at least 16 amino acids, preferably at least 20 amino acids, more preferably at least 25 amino acids, and most preferably 35 amino acids.

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For nucleic acids, the length of comparison sequences will generally be at least 50 nucleotides, preferably at least 60 nucleotides, more preferably at least 75 nucleotides, and most preferably 110 nucleotides.

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By "substantially identical" is meant an amino acid

sequence which differs only by conservative amino acid substitutions, for example, substitution of one amino acid for another of the same class (e.g., valine for glycine, arginine for lysine, etc.) or by one or more nonconservative substitutions, deletions, or insertions located at positions of the amino acid sequence which do not destroy the function of the protein (assayed, e.g., as described herein). Preferably, such a sequence is at least 85%, more preferably 90%, more preferably 95%, more preferably 98%, and most preferably 99% identical at the amino acid level to one of the sequences of EGFP (SEO ID NO:), EYFP (SEQ ID NO:), ECFP (SEQ ID NO:), EYFP-V68L/Q69K (SEQ ID NO: __), YFP H148G (SEQ ID NO:__), or YFP H148Q (SEQ ID NO:___) . Homology is typically measured using sequence analysis software (e.g., Sequence Analysis Software Package of the Genetics Computer Group, University of Wisconsin Biotechnology Center, 1710 University Avenue, Madison, WI 53705). Such software matches similar sequences by

assigning degrees of homology to various substitutions, deletions, substitutions, and other modifications. Conservative substitutions typically include substitutions within the following groups: glycine alanine; valine, isoleucine, leucine; aspartic acid, glutamic acid, apparagine, glutamine; serine, threonine; lysine, arginine;

and phenylalanine, tyrosine.

In some embodiments, the amino acid sequence of the protein includes one of the following sets of substitutions

in the amino acid sequence of the Aequora green fluorescent protein (SEQ ID NO:_): F64L/S65T/H231L, referred to herein as EGFP (SEQ ID NO:_); S65G/S72A/T203Y/H231L, referred to herein as EYFP (SEQ ID NO:_); S65G/V68L/Q69K/S72A/T203Y/H231L, referred to herein as EYFP-V68L/Q69K (SEQ ID NO:_); X26R/F64L/S65T/Y66W/N146I/M153T/V163A/N164H/H231L, referred to herein as ECFP (SEQ ID NO:_). The amino acid sequences of EGFP, EYFP, ECFP, and EYFP-V68L/Q69K are shown in Tables 1-4, respectively. The amino acids are numbered with the amino acid following the initiating methionine assigned the '1' position. Thus, F64L corresponds to a substitution of leucine for phenylalanine in the 64th amino acid following the initiating methionine.

Table 1. EGFP Amino Acid Sequence (SEQ ID NO:__)

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MVSKGEELFTGVVPILVELDGDVNGHKFSVSGEGEGDATYGKLTLKFICTTGKLPVPWPT LVTTLTTGVQCFSRYPDHNKOHDFFKSAMPEGTVORTIFFKDDGNYKTRABVKFEGDTL VNRIELKGIDFKEDGNILGHKLEYNYNSHNVYIMADKQKNGIKVNFKIRHNIEDGSVQLA DHYOONTPIGDGPVLLPDNHYLSTQSALSKDPNEKRDHMVLLEFVTAAGITLGMDELYK

Table 2. EYFP Amino Acid Sequence (SEQ ID NO:)

MVSKGEELFTGVVPILVELDGDVNGHKFSVSGEGEGDATYGKLTLKFICTTGKLPVPMPT LVTTFGYGVQCFARYPDHMKQHDFFKSAMPEGYVQERTIFFKDGNYKTRAEVKEEGIT VNRIELKGIDFKEDGNILGHKLEYNYNSHNVYIMADKQKNGIKVNFKIRHNIEDGSVQLA DHYQQNTPIGDGPVLLPDNHYLSYQSALSKDPNEKRDHMVLLEFVTAAGITIGMDELYK

Table 3. EYFP-V68L/Q69K Amino Acid Sequence (SEQ ID NO:__)

MVSKGEELFTGVVPILVELDGDVNGHKFSVSGEGEGDATYGKLTLKFICTTGKLPVPWPT
LVTTFGYGLKCFARYPDHMKQHDFFKSAMPEGTVQERTIFFKDDGNYKTRAEVKFEGDTL
VNRIELKGIDFKEDGNILGHKLEYNYNSHNVYIMADKQKNGIKVNFKIRHNIEDGSVQLA
DHYOONTPIGDGPVLLPDNHYLSVOSALSKDPMEKRDHWVLLEFVTAAGITIGMDELYK

Table 4. ECFP Amino Acid Sequence (SEQ ID NO:__)

MVSKGEELFTGVVPILVELDGDVNGHRFSVSGEGEGDATYGKLTLKFICTTGKLPVPWPT

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LVTTLTWGVQCFSRYPDHMKQHDFFKSAMPBGYVQBRTIFFKDDGNYKTRAEVKFBGDTL"
VNRIELKGIDFKBDGNILGHKUEYNYISHNYVITADKQKMGIKAHFKIRINIEDGSVQLA
DHYQONTPIOGGVULPDNHYLSTQSALSKDPNBKRDHWLLEFVTAAGITLGMDELYK

In other embodiments, the amino acid sequence of the protein is based on the sequence of the wild-type Aeguora green fluorescent protein, but includes the substitution H148G (SEQ ID NO:__) or H148Q (SEQ ID NO:__). In specific embodiments, these substitutions can be present along with other substitutions, e.g., the proteins can include the substitutions S65G/V68L/S72A/H148G/Q80R/T203Y (SEQ ID NO:__), which is referred to herein as the "YFP H148G mutant," S65G/V68L/S72A/H148Q/Q80R/T203Y, which is referred to herein as the "YFP H148G wutant" (SEQ ID NO:__), the as well as EYFP-H148G (SEQ ID NO:__) and EFP-H148Q (SEQ ID NO:__). The amino acid sequences of these mutants are shown in Tables 5-8, respectively.

Table 5. Amino Acid Sequence of YFP H148G (SEQ ID NO:__)

MSKGEELFTGVVPILVELDGDVNGHKFSVSGEGEGDATYGKLTLKPICTTGKLPVPWPTL
VTTFGYGLQCFARYPDHWRHDFFKSAMPEGYVQERTIFFKDDGNYKTRAEVKFEGDTLV
20 NRIELKGIDFKEDGNILGHKLEYNYNSGNVYIMADKQKNGIKVNFKIRHNIEDGSVQLAD
HYQQNTPIGDGPVLLPDNHYLSYQSALSKDPNEKRDHMVLLEFVTAAGITHGMDELYK

Table 6. Amino Acid Sequence of YFP H148Q (SEQ ID NO:__)
MSKGEBLFTGVVPILVELDGDVNGHKFSVSGEGEGDATYGKLTLKFICTTGKLPVPWPTL
VTTFGYGLQCFARYPDHMKRHDFFKSAMPEGYVQERTIFFKDDGNYKTRAEVKFBGDTLV
NR IBLKGIDFKEDGNILGHKLEYNYNSONVYIMADKÇKNGIKVNFKIRHNI EDGSVQLAD
HYQONTPIGDGPVLLPDNNYLSYSOALSKPDNEKDHWVLLEFVTAAGITHGMBELYK

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Table 7. Amino Acid Sequence of EYPP-H148G (SEQ ID NO:_)

MVSKGEELFTGVVPILVELDGDVMGHKFSVSGEGBGDATYGKLTLKFICTTGKLPVPWPT

LVTTFGGVQCFARYPDHMKCHDFPKSAMPEGYVQBRTIFFKDDGNYKTRABVKFEGDTL

VNRIELKGIDFKBDGNILGHKLBYNYNSGNVYIMADKQKNGIKVNFKIRHNIEDGSVQLA

DHYOONTPICHGPVLLPDNHYLSVGSALISKDPMEKERDHWVLLEFVTAAGTILGMDELYK

Table 8. Amino acid Sequence of EYFP-H148Q (SEQ ID NO:__)

MVSKGEELFTGVVPILVELDGDVNGHKFSVSGEGEGDATYGKLTLKFICTTGKLPVPWPT -LVTTFGYGVQCFARYEDHMKQHDFFKSAMPEGYVQERTIFFKDDGNYKTRAEVKFBGDTL VNRIELKGIDFKEDGNILGHKLEYNYNSQNVYIMADKQKNGIKVNFKIRHNIEDGSVQLA DHYQONTPIGDGPVLLPDNHYLSYQSALSKDPNEKRDHMVLLEFVTAAGITLGMDELYK

In some embodiments, the protein or polypeptide is substantially purified. By "substantially pure protein or polypeptide" is meant an functional engineered fluorescent polypeptide which has been separated from components which naturally accompany it. Typically, the protein or polypeptide is substantially pure when it is at least 60%, 10 by weight, free from the proteins and naturally-occurring organic molecules with which it is naturally associated. Preferably, the preparation is at least 75%, more preferably at least 90%, and most preferably at least 99%, by weight, of the protein. A substantially pure protein may be 15 obtained, for example, by extraction from a natural source (e.g., a plant cell); by expression of a recombinant nucleic acid encoding a functional engineered fluorescent protein; or by chemically synthesizing the protein. Purity can be measured by any appropriate method, e.g., those described in 20 column chromatography, polyacrylamide gel electrophoresis, or by HPLC analysis.

A protein or polypeptide is substantially free of naturally associated components when it is separated from 25 those contaminants which accompany it in its natural state. Thus, a protein or polypeptide which is chemically synthesized or produced in a cellular system different from the cell from which it naturally originates will be substantially free from its naturally associated components. Accordingly, substantially pure polypeptides include those derived from eukaryotic organisms but synthesized in E. coli or other prokaryotes.

The invention also provides polynucleotides encoding the functional engineered fluorescent protein described herein. These polynucleotides include DNA, cDNA, and RNA sequences which encode functional engineered fluorescent proteins. It is understood that all polynucleotides encoding functional engineered fluorescent proteins are also included herein, as long as they encode a protein or polypeptide whose fluorescent emission intensity changes as pH varies between 5 and 10. Such polynucleotides include naturally occurring, synthetic, and intentionally 1.0 manipulated polynucleotides. For example, the polynucleotide may be subjected to site-directed mutagenesis. The polynucleotides of the invention include sequences that are degenerate as a result of the genetic 15 code. Therefore, all degenerate nucleotide sequences are included in the invention as long as the amino acid sequence of the functional engineered fluorescent protein or derivative is functionally unchanged. Specifically disclosed herein is a polynucleotide sequence encoding a functional engineered fluorescent 20 protein that includes one of the following sets of substitutions in the amino acid sequence of the Aequora green fluorescent protein (SEQ ID NO:): S65G/S72A/T203Y/H231L, S65G/V68L/O69K/S72A/T203Y/H231L, or

protein that includes one of the following sets of substitutions in the amino acid sequence of the Aequora green fluorescent protein (SEQ ID NO:__):

S65G/S72A/T203Y/H231L, S65G/V68L/Q69K/S72A/T203Y/H231L, or

K26R/F64L/S65T/Y66W/N146I/M153T/V163A/N164H/H231L. In specific embodiments, the DNA sequences encoding EGFP, EYFP, ECFP, EYFP-V68L/Q69K, YFP H148G, and YFP H148Q are those shown in Table 9-16 (SEQ ID NOs: __ to __), respectively.

The nucleic acid encoding functional engineered

30 fluorescent proteins may be reflect the codon choice in the native A. victoria coding sequence, or, alternatively, may be chosen to reflect the optimal codon frequencies used in the organism in which the proteins will be expressed. Thus,

nucleic acids encoding a target functional engineered protein to be expressed in a human cell may have use a codon choice that is optimized for mammals, or especially humans.

Table 9. EGFP Nucleic Acid Sequence (SEQ ID NO:)

Table 10. EYFP Nucleic Acid Sequence (SEQ ID NO: __)

Table 11. ECFP Nucleic Acid Sequence (SEQ ID NO:)

GACCACTACCAGCAGAACACCCCCATCGGCGACGGCCCCGTGCTGCTGCCCGACAACCACTACCTGAGCACCCAGTCCGGCCTGAGCAAAGACCCCAACAGAAGCCGGATCACATGGTCCTGCTGAGTTCTTGGAGTTCTTGTGAGTTCTTGGAGTTCTTGGAGTTCTTGGAGTTCTTGGAGTTCTTGAGAGTCATTACAAGTAA

Table 12. EYFP-V68L/Q69K Nucleic Acid Sequence (SEO ID NO:)

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Table 13. Nucleotide Sequence of the YFP H148G Coding Region (SEQ ID NO:)

Table 14. Nucleotide Sequence of the YFP H148Q Coding Region

ATGAGTAAAGGACAACATTTTCACTGAGTTGTCCCAATTCTTGTTGAATTAGATGGT GATGTTAATTGGCACAAATTTTCTGTCAGTGGAGAGGGTGAAGGTGATGCAACATACGGA AAACTTACCCTTAAATTTATTTGCACTACTGGAAAACTACCTGGTTCCATGGCCAACACT GTCACTACTTTCGGTTATGGTCTTCAATGCTTTGCAAGATACCCAGATCATATGAAACGG CATGACTTTTTCAAGAGTGCCATGCCGGAAGGTTAGTTCAGGAAAGAACTATATTTTTC AAAGATGACGGGAACTACAAGACGTGCTGAAGTCAAGTTTGAAGGTGATACCCTTGTT AATGAATCAGGTTAAAAGGTATTGATTTTAAAGAAGATGGAAACATTCTTGGACACAAA

Table 15. Nucleotide Sequence of the EYFP-H148G Coding Region

20 Table 16. Nucleotide Sequence of the EYFP-H148Q Coding Region

The term "polynucleotide" refers to a polymeric form

of nucleotides of at least 10 bases in length. The
nucleotides can be ribonucleotides, deoxyribonucleotides, or
modified forms of either type of nucleotide. The term
includes single and double stranded forms of DNA. By

"isolated polynucleotide" is meant a polynucleotide that is not immediately contiguous with both of the coding sequences with which it is immediately contiguous (one on the 5' end and one on the 3' end) in the naturally occurring genome of the organism from which it is derived. The term therefore includes, for example, a recombinant DNA which is incorporated into a vector, e.g., an expression vector; into an autonomously replicating plasmid or virus; or into the genomic DNA of a prokaryote or eukaryote, or which exists as a separate molecule (e.g., a cDNA) independent of other sequences.

A "substantially identical" nucleic acid sequence codes for a substantially identical amino acid sequence as defined above.

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15 The functional engineered fluorescent protein can also include a targeting sequence to direct the fluorescent protein to particular cellular sites by fusion to appropriate organellar targeting signals or localized host proteins. A polynucleotide encoding a targeting sequence 20 can be ligated to the 5' terminus of a polynucleotide encoding the fluorescence such that the targeting peptide is located at the amino terminal end of the resulting fusion polynucleotide/polypeptide. The targeting sequence can be, e.g., a signal peptide. In the case of eukaryotes, the signal peptide is believed to function to transport the 25 fusion polypeptide across the endoplasmic reticulum. The secretory protein is then transported through the Golgi apparatus, into secretory vesicles and into the extracellular space or, preferably, the external 30 environment. Signal peptides which can be utilized according to the invention include pre-pro peptides which contain a proteolytic enzyme recognition site. Other signal peptides with similar properties to pro-calcitonin described

herein are known to those skilled in the art, or can be readily ascertained without undue experimentation. The targeting sequence can also be a nuclear localization sequence, an endoplasmic reticulum localization sequence, a peroxisome localization sequence, a mitochondrial localization sequence, or a localized protein. Targeting sequences can be targeting sequences which are described, for example, in "Protein Targeting", chapter 35 of Stryer, L., Biochemistry (4th ed.). W.H. Freeman, 1995. The localization sequence can also be a localized protein. 10 Some important targeting sequences include those targeting the nucleus (KKKRK) (SEQ ID NO:), mitochondrion (the 12 amino terminal amino acids of the cytochrome c oxidase subunit IV gene (SEQ ID NO:), or the amino terminal sequence MLRTSSLFTRRVQPSLFRNILRLQST (SEQ ID NO:), 15 endoplasmic reticulum (KDEL (SEO ID NO:) at the C-terminus, assuming a signal sequence present at N-terminus), peroxisome (SKF at C-terminus), prenylation or insertion into plasma membrane (CaaX, CC, CXC, or CCXX at C-terminus), cytoplasmic side of plasma membrane (fusion to SNAP-25), or the Golgi apparatus (fusion to the amino terminal 81 amino acids of human type II membrane-anchored

furin).

Examples of targeting sequences linked to functional engineered fluorescent proteins include GT-EYFP (SEQ ID NO:__), GT-EGFP (SEQ ID NO:__), and GT-EYFP-V68L/Q69K (SEQ ID NO:__), which are targeted to the Golgi apparatus using sequences from the GT protein; and

protein galactosyltransferase (SEO ID NO:), or fusion to

30 EYFP-mito (SEQ ID NO:__) and EGFP-mito (SEQ ID NO:__), which are targeted to the mitochondrial matrix using sequences from the amino terminal region of the cytochrome c oxidase subunit IV gene. The EYFP, EGFP, EGFP, and EYFP-V68L/069K

amino acid sequences, as well as nucleic acids encoding these polypeptides, are described above. The GT-derived targeting sequence corresponds to the 81 amino terminal amino acids of the human GT sequence. The GT amino acid sequences, and the polynucleotide sequences encoding the GT amino acid sequences, are described in Genbank Accession No. M70427 and Mengle-Gaw et al., Biochem. Biophys. Res. Commun. 176 (3), 1269-1276 (1991).

Amino acid sequences of mito-ECFP, mito-EYFP, GT
10 EGFP, and GT-EYFP, mito-YFP H148G, mito-YFP H148G, mito-EYFP

H148G, mito YFP-H148O are shown in Tables 17-24.

In specific embodiments, nucleic acid sequences encoding targeting sequences linked to functional engineered fluorescent proteins have the sequences shown in Tables 23-32.

Table 17. mito-ECFP Amino Acid Sequence (SEQ ID NO: __)

MLSLRQSIRFFKRSGIMVSKGEELFTGVVPILVELDGDVNGHRFSVSGEGEGDATYGKLT
LKFICTTGKLPVPMPTLVTTLTWGVQCFSRYPDHMKQHDFFKSAMPEGYYQERTIFFKDG
GNYKTRAEVKFEGDTLVNRIELKGIDFKEDGNILGHKLEYNYISHNVYLTADAQKNGIK
HFKIRRINI BDGSVULADPHYQONTPIEJGEPVLLPDHYLYSTOSALSKDPNEKRPHWYLLEF

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VTAAGITLGMDELYK

Table 18. mito-EYFP Amino Acid Sequence (SEQ ID NO:) MLSLROSIRFFKRSGIMVSKGEELFTGVVPILVELDGDVNGHKFSVSGEGEGDATYGKLT LKFICTTGKLPVPWPTLVTTFGYGVQCFARYPDHMKQHDFFKSAMPEGYVQERTIFFKDD GNYKTRAEVKFEGDTLVNRIELKGIDFKEDGNILGHKLEYNYNSHNVYIMADKOKNGIKV NFKIRHNIEDGSVQLADHYQONTPIGDGPVLLPDNHYLSYQSALSKDPNEKRDHMVLLEF VTAAGITLGMDELYK Table 19. GT-EGFP Amino Acid Sequence (SEQ ID NO:) MRLREPLLSGAAMPGASLORACRLLVAVCALHLGVTLVYYLAGRDLSRLPOLVGVSTPLO GGSNSAAATGOSSGELRTGGAMDPMVSKGEELFTGVVPILVELDGDVNGHKFSVSGEGEG DATYGKLTLKFI CTTGKLPVPWPTLVTTLTYGVOCFSRYPDHMKOHDFFKSAMPEGYVOE RTIFFKDDGNYKTRAEVKFEGDTLVNRIELKGIDFKEDGNILGHKLEYNYNSHNVYIMAD KOKNGI KVNFKI RHNI EDGSVOLADHYOONTPI GDGPVLLPDNHYLSTOSALSKDPNEKR DHMVLLEFVTAAGITLGMDELYK Table 20. GT-EYFP Amino Acid Sequence (SEQ ID NO:) MRLREPLLSGAAMPGASLQRACRLLVAVCALHLGVTLVYYLAGRDLSRLPQLVGVSTPLQ GGSNSAAAIGOSSGELRTGGAMDPMVSKGEELFTGVVPILVELDGDVNGHKFSVSGEGEG DATYGKLTLKFICTTGKLPVPWPTLVTTFGYGVQCFARYPDHMKQHDFFKSAMPEGYVOE RTIFFKDDGNYKTRAEVKFEGDTLVNRIELKGIDFKEDGNILGHKLEYNYNSHNVYIMAD KOKNGIKVNFKIRHNIEDGSVOLADHYOONTPIGDGPVLLPDNHYLSYOSALSKDPNEKR DHMVLLEFVTAAGITLGMDELYK* Table 21. mito-YFP-H148G Amino Acid Sequence (SEO ID NO:) MLRTSSLFTRRVOPSLFRNILRLOSTSKGEELFTGVVPILVELDGDVNGHKFSVSGEGEG DATYGKLTLKFICTTGKLPVPWPTLVTTFGYGLQCFARYPDHMKRHDFFKSAMPEGYVQE RTIFFKDDGNYKTRAEVKFEGDTLVNRIELKGIDFKEDGNILGHKLEYNYNSGNVYIMAD KOKNGTKVNFKTRHNTEDGSVOLADHYOONTPIGDGPVLLPDNHYLSYOSALSKDPNEKR DHMVIJERFVTAAGTTHGMDRLYK

30 Table 22. mito-EYFP-H148Q Amino Acid Sequence (SEQ ID NO:_)
MRTSSLFTRRVQPSLFRNILRLQSTSKGEELFTGVVPILVELDGDVNGHKFSVSGEGEG
DATYGKLTLKFICTTGKLPVPWPTLVTTFGYGLQCFARYPDHMKRHDFFKSAMPEGYVQE
RTIFFKDGDNYKTRAEVKFEGDTLVNRIELKGIDFKEDGNILGHKLEYNNNSQNYVIMAD
KOKNGIKVNFKIRINIEDGSVOLADHYQONTPIODGPVLLIPDNYYLSYOSALSKDPNEKR

35 DHMVLLEFVTAAGITHGMDELYK

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Table 23. mito-EYFP-H148G Amino Acid Sequence (SEO ID NO:)

MLRTSSLFTRRVQPSLFRNILRLQSTMVSKGEELFTGVVPILVELDGDVNGHKFSVSGEG EGDATYGKLTLKFICTTGKLPVPMPTLVTTEGYGVQCPARYPDHMKQHDFFKSAMPEGVV QERTIFFKDDGNYKTRAEVKFEGDTLVNRIELKGIDFKEDGNILGHKLEYNYNSGNVYIM ADKQRNGIKVNFRIRHNIEDGSVQLADHYQQNTPIGDGPVLLPDNHYLSYQSALSKDPNE KRDHWYLLEFVTAAGTILGMDELYK

Table 24. mito-EYFP-H148Q Amino Acid Sequence (SEQ ID NO:)

- 10 MLRTSSLETERVQPSLERNILKLQSTMYSKGEBLEFTGYVPILVELDGDVNGHKFSVSGEG EGDATYGKLTLKFICTTGKLPVFWFTLVTTFGYGVQCFARYPDHMKQHDFFKSAMPEGYV QERTIFFKDDGNYKTRAEVKFEGDTLVNRIELKGIDFKEDGNILGHKLEVNYNSQNVYIM ADKQKNGIKVNFKIRNNIEDGSVQLADHYQQNTPIGDGPVLLPDNHYLSYQSALSKDPNE KRDHVYLLEFVTAGATILKMDELIK
- 15 Table 25. GT-ECFP Nucleic Acid Sequence (SEQ ID NO: __)

ARGAGCTTCGGGAGCGCTCCTGAGCGGCGCGCGATGCCAGGGGGTCCCTACAGCGG GCCTGCCGCTGTGGGGCTCTGGGGTCTTGGGTCACCTTGGGTCACCCTGTTTACTAC CTGGCTGGCGCGACCTGAGCGCCTGCCCCAACTGGTGGAGTTCCACACCGCTGCAG GGCGCTCGAACAGTGCCGCCATCGGGCAGTCCTCCGGGGACCTCCGGACCGACGAGGG 20 GCCATGGATCCCATGCTAGACAGGGCGAGGAGCTTCACCGGGGTGGTGCCCATCCTG GTCGAGCTGGACGGCGACGTAAACGGCCACAGGTTCACGGTGTCCCGGCGAGGGCGAGGGC

- GTGCAGCTCGCCGACCACTACCAGCAGAACACCCCCATCGGCGACGGCCCCGTGCTGCTG
 CCCGACAACCACTACCTGAGCACCCAGTCCGCCCTGAGCAAAGACCCCAACGAGAAGGGC
 GATCACATGGTCCTGCTGGAGTTCGTGACCGCCGCGGGATCACTCTCGGCATGGACGAG
 CTGTACAAGTAA

Table 26. mito-EYFP Nucleic Acid Sequence (SEQ ID NO: __)

Table 27. GT-EGFP Nucleic Acid Sequence (SEQ ID NO:__)

10 ATGAGGCTTCGGGAGCCGCTCCTGAGCGGCGCGCGCGATGCCAGGCGCGTCCCTACAGCGG GCCTGCCGCCTGCTCGTGGCCGTCTGCGCTCTGCACCTTGGCGTCACCCTCGTTTACTAC CTGGCTGGCCGCGACCTGAGCCGCCTGCCCCAACTGGTCGGAGTCTCCACACCGCTGCAG GGCGGCTCGAACAGTGCCGCCGCCATCGGGCAGTCCTCCGGGGAGCTCCGGACCGGAGGG 15 GCCATGGATCCCATGGTGAGCAAGGGCGAGGAGCTGTTCACCGGGGTGGTGCCCATCCTG GTCGAGCTGGACGCGACGTAAACGGCCACAAGTTCAGCGTGTCCGGCGAGGGCGAGGGC GATGCCACCTACGGCAAGCTGACCCTGAAGTTCATCTGCACCACCGGCAAGCTGCCCGTG CCCTGGCCCACCCTCGTGACCACCCTGACCTACGGCGTGCAGTGCTTCAGCCGCTACCCC GACCACATGAAGCAGCACGACTTCTTCAAGTCCGCCATGCCCGAAGGCTACGTCCAGGAG CGCACCATCTTCTTCAAGGACGACGGCAACTACAAGACCCGCGCGGAGGTGAAGTTCGAG 20 GGCGACACCCTGGTGAACCGCATCGAGCTGAAGGGCATCGACTTCAAGGAGGACGGCAAC ATCCTGGGGCACAAGCTGGAGTACAACTACAACAGCCACAACGTCTATATCATGGCCGAC AAGCAGAAGAACGGCATCAAGGTGAACTTCAAGATCCGCCACAACATCGAGGACGGCAGC GTGCAGCTCGCCGACCACTACCAGCAGAACACCCCCATCGGCGACGGCCCCGTGCTGCTG CCCGACACCACTACCTGAGCACCCAGTCCGCCCTGAGCAAAGACCCCAACGAGAAGCGC 25 GATCACATGGTCCTGCTGGAGTTCGTGACCGCCGCGGGATCACTCTCGGCATGGACGAG CTGTACAAGTAA

Table 28. GT-EYFP Nucleic Acid Sequence (SEO ID NO:)

ATGAGGCTTCGGGAGCCGCTCCTGAGCGGCGCCGCGATGCCAGGCGCGTCCCTACAGCGG 30 GCCTGCCGCCTGCTCGTGGCCGTCTGCGCTCTGCACCTTGGCGTCACCCTCGTTTACTAC CTGGCTGGCCGCGCCTGAGCCGCCTGCCCCAACTGGTCGGAGTCTCCACACCGCTGCAG GGCGGCTCGAACAGTGCCGCCGCCATCGGGCAGTCCTCCGGGGGGGCTCCGGACCGGAGGG GCCATGGATCCCATGGTGAGCAAGGGCGAGGAGCTGTTCACCGGGGTGGTGCCCATCCTG GTCGAGCTGGACGCGACGTAAACGGCCACAAGTTCAGCGTGTCCGGCGAGGGCGAGGGC 35 GATGCCACCTACGGCAAGCTGACCCTGAAGTTCATCTGCACCACCGGCAAGCTGCCCGTG CCCTGGCCCACCTCGTGACCACCTTCGGCTACGGCTGCAGTGCTTCGCCCGCTACCCC GACCACATGAAGCACGACTTCTTCAAGTCCGCCATGCCCGAAGGCTACGTCCAGGAG CGCACCATCTTCTTCAAGGACGACGCCAACTACAAGACCCGCGCGAGGTGAAGTTCGAG GGCGACACCCTGGTGAACCGCATCGAGCTGAAGGGCATCGACTTCAAGGAGGACGGCAAC ATCCTGGGGCACAAGCTGGAGTACAACTACAACAGCCACAACGTCTATATCATGGCCGAC 40 AAGCAGAAGAACGGCATCAAGGTGAACTTCAAGATCCGCCACAACATCGAGGACGGCAGC GTGCAGCTCGCCGACCACTACCAGCAGAACACCCCCATCGGCGACGGCCCCGTGCTGCTG CCCGACACCACTACCTGAGCTACCAGTCCGCCCTGAGCAAAGACCCCAACGAGAAGCGC GATCACATGGTCCTGCTGGAGTTCGTGACCGCCGCCGGGATCACTCTCGGCATGGACGAG 45 CTGTACAAGTAA

Table 29. mito-YFP H148G Nucleic Acid Sequence (SEQ ID NO:)

Table 30. mito-YFP H148Q Nucleic Acid Sequence (SEQ ID NO:)

ATGCTGAGCCTGCGCCAGAGCATCCGCTTCTTCAAGCGCAGCGGCATCATGAGTAAAGGA GAAGAACTTTTCACTGGAGTTGTCCCAATTCTTGTTGAATTAGATGGTGATGTTAATGGG 20 CACAAATTTTCTGTCAGTGGAGAGGGTGAAGGTGATGCAACATACGGAAAACTTACCCTT AAATTTATTTGCACTACTGGAAAACTACCTGTTCCATGGCCAACACTTGTCACTACTTTC GGTTATGGTCTTCAATGCTTTGCAAGATACCCAGATCATATGAAACGGCATGACTTTTTC AAGAGTGCCATGCCCGAAGGTTATGTTCAGGAAAGAACTATATTTTTCAAAGATGACGGG AACTACAAGACACGTGCTGAAGTCAAGTTTGAAGGTGATACCCTTGTTAATAGAATCGAG TTAAAAGGTATTGATTTTAAAGAAGATGGAAACATTCTTGGACACAAATTGGAATACAAC 25 TTCAAAATTAGACACACATTGAAGATGGAAGCGTTCAACTAGCAGACCATTATCAACAA AATACTCCAATTGGCGATGGCCCTGTCCTTTTACCAGACAACCATTACCTGTCCTATCAA TCTGCCCTTTCGAAAGATCCCAACGAAAAGAGAGACCACATGGTCCTTCTTGAGTTTGTA 30 ACAGCTGCTGGGATTACACATGGCATGGATGAACTATACAAA

Table 31. mito-EYFP-H148G Nucleic Acid Sequence (SEQ ID NO:

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Table 32. mito-EYFP-H148Q Nucleic Acid Sequence (SEQ ID NO:)

ATGCTGAGCCTGCGCCAGAGCATCCGCTTCTTCAAGCGCAGCGCATCATGGTGAGCAAG GGCGAGGAGCTGTTCACCGGGGTGGTGCCCATCCTGGTCGAGCTGGACGGCGACGTAAAC GGCCACAGTTCAGCGTGTCCGGCGAGGGCGAGGGCGATGCCACCTACGGCAAGCTGACC 20 CTGAAGTTCATCTGCACCACCGGCAAGCTGCCCGTGCCCTGGCCCACCCTCGTGACCACC TTCGGCTACGGCGTGCACTGCTCCCCGCTACCCCGACCACATGAAGCACGACTTC TTCAAGTCCGCCATGCCCGAAGGCTACGTCCAGGAGCGCACCATCTTCTTCAAGGACGAC GGCAACTACAAGACCCGCGCCGAGGTGAAGTTCGAGGGCGACACCCTGGTGAACCGCATC 25 GAGCTGAAGGGCATCGACTTCAAGGAGGACGGCAACATCCTGGGGCACAAGCTGGAGTAC AACTACAACAGCCAGAACGTCTATATCATGGCCGACAAGCAGAAGAACGGCATCAAGGTG AACTTCAAGATCCGCCACAACATCGAGGACGGCAGCGTGCAGCTCGCCGACCACTACCAG CAGAACACCCCCATCGGCGACGGCCCCGTGCTGCTGCCCGACAACCACTACCTGAGCTAC CAGTCCGCCTGAGCAAAGACCCCAACGAGAAGCGCGATCACATGGTCCTGCTGGAGTTC 3.0 GTGA CCGCCGCGGGATCACTCTCGGCATGGACGAGCTGTACAAGTAA

The fluorescent indicators can be produced as proteins fused to other fluorescent indicators or targeting sequences by recombinant DNA technology. Recombinant production of fluorescent proteins involves expressing nucleic acids having sequences that encode the proteins. Nucleic acids encoding fluorescent proteins can be obtained by methods known in the art. For example, a nucleic acid encoding the protein can be isolated by polymerase chain reaction of cDNA from A. victoria using primers based on the

DNA sequence of A. victoria green fluorescent protein. PCR methods are described in, for example, U.S. Pat. No. 4,683,195; Mullis, et al. Cold Spring Harbor Symp. Quant. Biol. 51:263 (1987), and Erlich, ed., PCR Technology, (Stockton Press, NY, 1989). Mutant versions of fluorescent proteins can be made by site-specific mutagenesis of other nucleic acids encoding fluorescent proteins, or by random mutagenesis caused by increasing the error rate of PCR of the original polynucleotide with 0.1 mM MnCl₂ and unbalanced nucleotide concentrations. See, e.g., U.S. patent application 08/337,915, filed November 10, 1994 or International application PCT/US95/14692, filed 11/10/95.

The construction of expression vectors and the

expression of genes in transfected cells involves the use of
molecular cloning techniques also well known in the art.
Sambrook et al., Molecular Cloning -- A Laboratory Manual,
Cold Spring Harbor Laboratory, Cold Spring Harbor, NY,
(1989) and Current Protocols in Molecular Biology, F.M.
Ausubel et al., eds., (Current Protocols, a joint venture
between Greene Publishing Associates, Inc. and John Wiley &
Sons, Inc., most recent Supplement).

Nucleic acids used to transfect cells with sequences coding for expression of the polypeptide of interest generally will be in the form of an expression vector

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including expression control sequences operatively linked to a nucleotide sequence coding for expression of the polypeptide. As used herein, "operatively linked" refers to a juxtaposition wherein the components so described are in a relationship permitting them to function in their intended manner. A control sequence operatively linked to a coding sequence is ligated such that expression of the coding sequence is achieved under conditions compatible with the control sequences. "Control sequence" refers to

polynucleotide sequences which are necessary to effect the expression of coding and non-coding sequences to which they are ligated. Control sequences generally include promoter, ribosomal binding site, and transcription termination sequence. The term "control sequences" is intended to include, at a minimum, components whose presence can influence expression, and can also include additional components whose presence is advantageous, for example, leader sequences and fusion partner sequences.

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As used herein, the term "nucleotide sequence coding for expression of" a polypeptide refers to a sequence that, upon transcription and translation of mRNA, produces the polypeptide. This can include sequences containing, e.g., introns. As used herein, the term "expression control sequences" refers to nucleic acid sequences that regulate the expression of a nucleic acid sequence to which it is operatively linked. Expression control sequences are operatively linked to a nucleic acid sequence when the expression control sequences control and regulate the transcription and, as appropriate, translation of the nucleic acid sequence. Thus, expression control sequences can include appropriate promoters, enhancers, transcription terminators, a start codon (i.e., ATG) in front of a protein-encoding gene, splicing signals for introns, maintenance of the correct reading frame of that gene to permit proper translation of the mRNA, and stop codons.

Methods which are well known to those skilled in the art can be used to construct expression vectors containing the fluorescent indicator coding sequence and appropriate transcriptional/translational control signals. These methods include in vitro recombinant DNA techniques, synthetic techniques and in vivo recombination/genetic recombination. (See, for example, the techniques described

in Maniatis, et al., Molecular Cloning A Laboratory Manual, Cold Spring Harbor Laboratory, N.Y., 1989). Transformation of a host cell with recombinant DNA may be carried out by conventional techniques as are well known to those skilled in the art. Where the host is prokaryotic, such as E. coli, competent cells which are capable of DNA uptake can be prepared from cells harvested after exponential growth phase and subsequently treated by the CaCl, method by procedures well known in the art. Alternatively, MgCl₂ or RbCl can be used. Transformation can also be performed after forming a protoplast of the host cell or by electroporation.

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When the host is a eukaryote, such methods of transfection of DNA as calcium phosphate co-precipitates, conventional mechanical procedures such as microinjection, electroporation, insertion of a plasmid encased in liposomes, or virus vectors may be used. Eukaryotic cells can also be cotransfected with DNA sequences encoding the fusion polypeptide of the invention, and a second foreign DNA molecule encoding a selectable phenotype, such as the herpes simplex thymidine kinase gene. Another method is to use a eukaryotic viral vector, such as simian virus 40 (SV40) or bovine papilloma virus, to transiently infect or transform eukaryotic cells and express the protein. (Eukaryotic Viral Vectors, Cold Spring Harbor Laboratory, Gluzman ed., 1982).

Techniques for the isolation and purification of polypeptides of the invention expressed in prokaryotes or eukaryotes may be by any conventional means such as, for example, preparative chromatographic separations and immunological separations such as those involving the use of monoclonal or polyclonal antibodies or antiqen.

A variety of host-expression vector systems may be utilized to express fluorescent indicator coding sequence.

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These include but are not limited to microorganisms such as bacteria transformed with recombinant bacteriophage DNA, plasmid DNA or cosmid DNA expression vectors containing a fluorescent indicator coding sequence; yeast transformed 5 with recombinant yeast expression vectors containing the fluorescent indicator coding sequence; plant cell systems infected with recombinant virus expression vectors (e.g., cauliflower mosaic virus, CaMV; tobacco mosaic virus, TMV) or transformed with recombinant plasmid expression vectors (e.g., Ti plasmid) containing a fluorescent indicator coding sequence; insect cell systems infected with recombinant virus expression vectors (e.g., baculovirus) containing a fluorescent indicator coding sequence; or animal cell systems infected with recombinant virus expression vectors (e.g., retroviruses, adenovirus, vaccinia virus) containing 15 a fluorescent indicator coding sequence, or transformed animal cell systems engineered for stable expression. Depending on the host/vector system utilized, any of a number of suitable transcription and translation elements, including constitutive and inducible promoters, transcription enhancer elements, transcription terminators,

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20 etc. may be used in the expression vector (see, e.g., Bitter, et al., Methods in Enzymology 153:516-544, 1987). For example, when cloning in bacterial systems, inducible promoters such as pL of bacteriophage \(\lambda \), plac, ptrp, ptac 25 (ptrp-lac hybrid promoter) and the like may be used. When cloning in mammalian cell systems, promoters derived from the genome of mammalian cells (e.g., metallothionein promoter) or from mammalian viruses (e.g., the retrovirus long terminal repeat; the adenovirus late promoter; the 30 vaccinia virus 7.5K promoter) may be used. Promoters produced by recombinant DNA or synthetic techniques may also

be used to provide for transcription of the inserted fluorescent indicator coding sequence.

In bacterial systems a number of expression vectors may be advantageously selected depending upon the use intended for the fluorescent indicator expressed. For example, when large quantities of the fluorescent indicator are to be produced, vectors which direct the expression of high levels of fusion protein products that are readily purified may be desirable. Those which are engineered to contain a cleavage site to aid in recovering fluorescent 10 indicator are preferred. In yeast, a number of vectors containing constitutive or inducible promoters may be used. For a review see, Current Protocols in Molecular Biology, Vol. 2. Ed. Ausubel, et al., Greene Publish. Assoc. & Wiley Interscience, Ch. 13, 1988; Grant, et al., Expression and 15 Secretion Vectors for Yeast, in Methods in Enzymology, Eds. Wu & Grossman, 31987, Acad. Press, N.Y., Vol. 153, pp.516-544, 1987; Glover, DNA Cloning, Vol. II, IRL Press, Wash., D.C., Ch. 3, 1986; and Bitter, Heterologous Gene Expression in Yeast, Methods in Enzymology, Eds. Berger & 20 Kimmel, Acad. Press, N.Y., Vol. 152, pp. 673-684, 1987; and The Molecular Biology of the Yeast Saccharomyces, Eds. Strathern et al., Cold Spring Harbor Press, Vols. I and II, 1982. A constitutive yeast promoter such as ADH or LEU2 or an inducible promoter such as GAL may be used (Cloning in 25 Yeast, Ch. 3, R. Rothstein In: DNA Cloning Vol.11, A Practical Approach, Ed. DM Glover, IRL Press, Wash., D.C., 1986). Alternatively, vectors may be used which promote integration of foreign DNA sequences into the yeast

In cases where plant expression vectors are used, the expression of a fluorescent indicator coding sequence may be driven by any of a number of promoters. For example,

chromosome.

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viral promoters such as the 35S RNA and 19S RNA promoters of CaMV (Brisson, et al., Nature 310:511-514, 1984), or the coat protein promoter to TMV (Takamatsu, et al., EMBO J. 6:307-311, 1987) may be used; alternatively, plant promoters such as the small subunit of RUBISCO (Coruzzi, et al., 1984, EMBO J. 3:1671-1680: Broglie, et al., Science 224:838-843, 1984); or heat shock promoters, e.g., soybean hspl7.5-E or hsp17.3-B (Gurley, et al., Mol. Cell. Biol. 6:559-565, 1986) may be used. These constructs can be introduced into plant cells using Ti plasmids, Ri plasmids, plant virus vectors, 10 direct DNA transformation, microinjection, electroporation, etc. For reviews of such techniques see, for example, Weissbach & Weissbach, Methods for Plant Molecular Biology, Academic Press, NY, Section VIII, pp. 421-463, 1988; and Grierson & Corey, Plant Molecular Biology, 2d Ed., Blackie, 15 London, Ch. 7-9, 1988.

An alternative expression system which could be used to express fluorescent indicator is an insect system. In one such system, Autographa californica nuclear polyhedrosis virus (AcNPV) is used as a vector to express foreign genes. The virus grows in Spodoptera frugiperda cells. The fluorescent indicator coding sequence may be cloned into non-essential regions (for example, the polyhedrin gene) of the virus and placed under control of an AcNPV promoter (for example the polyhedrin promoter). Successful insertion of 25 the fluorescent indicator coding sequence will result in inactivation of the polyhedrin gene and production of non-occluded recombinant virus (i.e., virus lacking the proteinaceous coat coded for by the polyhedrin gene). These recombinant viruses are then used to infect Spodoptera 30 frugiperda cells in which the inserted gene is expressed, see Smith, et al., J. Viol. 46:584, 1983; Smith, U.S. Patent No. 4,215,051.

Eukaryotic systems, and preferably mammalian expression systems, allow for proper post-translational modifications of expressed mammalian proteins to occur. Eukaryotic cells which possess the cellular machinery for proper processing of the primary transcript, glycosylation, phosphorylation, and, advantageously secretion of the gene product should be used as host cells for the expression of fluorescent indicator. Such host cell lines may include but are not limited to CHO, VERO, BHK, HeLa, COS, MDCK, Jurkat, HEK-293, and WI38. Primary cell lines, such as neonatal rat myocytes, can also be used.

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Mammalian cell systems which utilize recombinant viruses or viral elements to direct expression may be engineered. For example, when using adenovirus expression vectors, the fluorescent indicator coding sequence may be ligated to an adenovirus transcription/translation control complex, e.g., the late promoter and tripartite leader sequence. This chimeric gene may then be inserted in the adenovirus genome by in vitro or in vivo recombination. Insertion in a non-essential region of the viral genome (e.g., region E1 or E3) will result in a recombinant virus that is viable and capable of expressing the fluorescent indicator in infected hosts (e.g., see Logan & Shenk, Proc. Natl. Acad. Sci. USA, 81: 3655-3659, 1984). Alternatively, the vaccinia virus 7.5K promoter may be used (e.g., see, Mackett, et al., Proc. Natl. Acad. Sci. USA ,79: 7415-7419, 1982: Mackett, et al., J. Virol. 49: 857-864, 1984; Panicali, et al., Proc. Natl. Acad. Sci. USA 79: 4927-4931, 1982). Of particular interest are vectors based on bovine papilloma virus which have the ability to replicate as extrachromosomal elements (Sarver, et al., Mol. Cell. Biol. 1: 486, 1981). Shortly after entry of this DNA into mouse cells, the plasmid replicates to about 100 to 200 copies per

cell. Transcription of the inserted cDNA does not require integration of the plasmid into the host's chromosome, thereby yielding a high level of expression. These vectors can be used for stable expression by including a selectable marker in the plasmid, such as the neo gene. Alternatively, the retroviral genome can be modified for use as a vector capable of introducing and directing the expression of the fluorescent indicator gene in host cells (Cone & Mulligan, Proc. Natl. Acad. Sci. USA, 81:6349-6353, 1984). High level expression may also be achieved using inducible promoters, including, but not limited to, the metallothionein IIA promoter and heat shock promoters.

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The recombinant nucleic acid can be incorporated into an expression vector including expression control sequences operatively linked to the recombinant nucleic acid. The expression vector can be adapted for function in prokaryotes or eukaryotes by inclusion of appropriate promoters, replication sequences, markers, etc.

DNA sequences encoding the fluorescence indicator
polypeptide of the invention can be expressed in vitro or in
vivo by DNA transfer into a suitable recombinant host cell.
As used herein, "recombinant host cells" are cells in which
a vector can be propagated and its DNA expressed. The term
also includes any progeny of the subject host cell. It is
understood that all progeny may not be identical to the
parental cell since there may be mutations that occur during
replication. However, such progeny are included when the
term "recombinant host cell" is used. Methods of stable
transfer, in other words when the foreign DNA is
continuously maintained in the host, are known in the art.

The expression vector can be transfected into a host cell for expression of the recombinant nucleic acid.

Recombinant host cells can be selected for high levels of

expression in order to purify the fluorescent indicator fusion protein. *E. coli* is useful for this purpose. Alternatively, the host cell can be a prokaryotic or eukaryotic cell selected to study the activity of an enzyme produced by the cell. In this case, the linker peptide is selected to include an amino acid sequence recognized by the protease. The cell can be, e.g., a cultured cell or a cell taken in vivo from a transgenic animal.

TRANSGENIC ANIMALS

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In another embodiment, the invention provides a transgenic non-human animal that expresses a polynucleotide sequence which encodes a fluorescent protein pH sensor.

The "non-human animals" of the invention comprise any non-human animal having a polynucleotide sequence which encodes a fluorescent indicator. Such non-human animals include vertebrates such as rodents, non-human primates, sheep, dog, cow, pig, amphibians, and reptiles. Preferred non-human animals are selected from the rodent family including rat and mouse, most preferably mouse. The "transgenic non-human animals" of the invention are produced by introducing "transgenes" into the germline of the non-human animal. Embryonal target cells at various developmental stages can be used to introduce transgenes. Different methods are used depending on the stage of development of the embryonal target cell. The zygote is the best target for micro-injection. In the mouse, the male pronucleus reaches the size of approximately 20 micrometers in diameter which allows reproducible injection of 1-2 pl of

DNA solution. The use of zygotes as a target for gene transfer has a major advantage in that in most cases the injected DNA will be incorporated into the host gene before the first cleavage (Brinster et al., Proc. Natl. Acad. Sci.

USA 82:4438-4442, 1985). As a consequence, all cells of the transgenic non-human animal will carry the incorporated transgene. This will in general also be reflected in the efficient transmission of the transgene to offspring of the founder since 50% of the germ cells will harbor the transgene. Microinjection of zygotes is the preferred method for incorporating transgenes in practicing the invention.

The term "transgenic" is used to describe an animal which includes exogenous genetic material within all of its cells. A "transgenic" animal can be produced by cross-breeding two chimeric animals which include exogenous genetic material within cells used in reproduction.

Twenty-five percent of the resulting offspring will be transgenic, i.e., animals which include the exogenous genetic material within all of their cells in both alleles.

50% of the resulting animals will include the exogenous

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genetic material within one allele and 25% will include no exogenous genetic material.

Retroviral infection can also be used to introduce transgene into a non-human animal. The developing non-human

embryo can be cultured in vitro to the blastocyst stage.

During this time, the blastomeres can be targets for retro viral infection (Jaenisch, R., Proc. Natl. Acad. Sci USA 73:1260-1264, 1976). Efficient infection of the blastomeres is obtained by enzymatic treatment to remove the zona pellucida (Hogan, et al. (1986) in Manipulating the Mouse Embryo, Cold Spring Harbor Laboratory Press, Cold Spring Harbor, N.Y.). The viral vector system used to introduce the transgene is typically a replication-defective retrovirus carrying the transgene (Jahner, et al., Proc. Natl. Acad. Sci. USA 82:6927-6931, 1985; Van der Putten, et al., Proc. Natl. Acad. Sci USA 82:6148-6152, 1985). Transfection is easily and efficiently obtained by culturing the blastomeres

on a monolayer of virus-producing cells (Van der Putten, supra; Stewart, et al., EMBO J. 6:383-388, 1987). Alternatively, infection can be performed at a later stage. Virus or virus-producing cells can be injected into the blastocoele (D. Jahner et al., Nature 298:623-628, 1982). Most of the founders will be mosaic for the transgene since incorporation occurs only in a subset of the cells which formed the transgenic nonhuman animal. Further, the founder may contain various retro viral insertions of the transgene at different positions in the genome which 10 generally will segregate in the offspring. In addition, it is also possible to introduce transgenes into the germ line, albeit with low efficiency, by intrauterine retro viral infection of the midgestation embryo (D. Jahner et al., 15 supra). A third type of target cell for transgene introduction is the embryonal stem cell (ES). ES cells are obtained from pre-implantation embryos cultured in vitro and fused with embryos (M. J. Evans et al. Nature 292:154-156, 1981; M.O. Bradley et al., Nature 309: 255-258,1984; Gossler, et al., Proc. Natl. Acad. Sci USA 83: 20 9065-9069, 1986; and Robertson et al., Nature 322:445-448, 1986). Transgenes can be efficiently introduced into the ES cells by DNA transfection or by retrovirus-mediated transduction. Such transformed ES cells can thereafter be combined with blastocysts from a nonhuman animal. The ES 25 cells thereafter colonize the embryo and contribute to the germ line of the resulting chimeric animal. (For review see Jaenisch, R., Science 240: 1468-1474, 1988).

"Transformed" means a cell into which (or into an ancestor of which) has been introduced, by means of recombinant nucleic acid techniques, a heterologous polynucleotide. "Heterologous" refers to a polynucleotide sequence that either originates from another species or is

modified from either its original form or the form primarily expressed in the cell.

"Transgene" means any piece of DNA which is inserted by artifice into a cell, and becomes part of the genome of the organism (i.e., either stably integrated or as a stable extrachromosomal element) which develops from that cell. Such a transgene may include a gene which is partly or entirely heterologous (i.e., foreign) to the transgenic organism, or may represent a gene homologous to an endogenous gene of the organism. Included within this 10 definition is a transgene created by the providing of an RNA sequence which is transcribed into DNA and then incorporated into the genome. The transgenes of the invention include DNA sequences which encode the fluorescent indicator which may be expressed in a transgenic non-human animal. The term 15 "transgenic" as used herein additionally includes any organism whose genome has been altered by in vitro manipulation of the early embryo or fertilized egg or by any transgenic technology to induce a specific gene knockout. The term "gene knockout" as used herein, refers to the 20 targeted disruption of a gene in vivo with complete loss of function that has been achieved by any transgenic technology familiar to those in the art. In one embodiment, transgenic animals having gene knockouts are those in which the target gene has been rendered nonfunctional by an insertion targeted to the gene to be rendered non-functional by homologous recombination. As used herein, the term "transgenic" includes any transgenic technology familiar to those in the art which can produce an organism carrying an

introduced transgene or one in which an endogenous gene has been rendered non-functional or "knocked out."

DETECTION OF DH USING FLUORESCENT INDICATOR PROTEINS

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In another embodiment, the invention provides a method for determining the pH of a sample by contacting the sample with an indicator including a first fluorescent protein moiety whose emission intensity changes as pH varies between pH 5 and 10, exciting the indicator, and then determining the intensity of light emitted by the first fluorescent protein moiety at a first wavelength. The emission intensity of the first fluorescent protein moiety indicates the pH of the sample.

The fluorescent protein moiety can be a functional engineered protein substantially identical to the amino acid sequence of Aequora green fluorescence protein (SEQ ID NO:-__). Preferred green fluorescence proteins include those having a functional engineered fluorescent protein that includes one of the following sets of substitutions in the amino acid sequence of the Aequora green fluorescent protein

S65G/V68L/Q69K/S72A/T203Y, or

Z66K/F64L/S65T/Y66W/N1461/M153T/V163A/N164H/H231L. Other preferred green fluorescence proteins include those having a functional engineered fluorescent protein that includes H148G or H148G substitutions in the Aequora green fluorescent protein. These proteins include the YFP H148G

25 (SEQ ID NO: __) and YFP H148Q (SEQ ID NO: __) proteins

(SEO ID NO:): S65G/S72A/T203Y/H231L,

described above.

The sample in which pH is to be measured can be a biological sample, e.g., a biological tissue such as an extracellular matrix, blood or lymphatic tissue, or a cell. The method is particularly suitable for measuring pH in a specific region of the cell, e.g., the cytosol, or an organellar space such as the inner mitochondrial matrix, the lumen of the Golqi, cytosol, the endoplasmic reticulum, the

chloroplast lumen, the lumen of lysosome, or the lumen of an endosome.

In some embodiments, the first fluorescent protein moiety is linked to a targeting sequence that directs the fluorescent protein to a desired cellular compartment. Examples of targeting sequences include the amino terminal 81 amino acids of human type II membrane-anchored protein galactosyltransferase (SEO ID NO:) for directing the fluorescent indicator protein to the Golgi and the amino terminal 12 amino acids of the presequence of subunit IV of 10 cytochrome c oxidase (SEQ ID NO:) for directing a fluorescent pH indicator protein to the mitochondrial matrix. The 12 amino acids of the presequence of subunit IV of cytochrome c oxidase (SEQ ID NO:) may be linked to the pH fluorescent indicator protein through a linker sequence, 15 e.g., Arg-Ser-Gly-Ile (SEQ ID NO:).

In another embodiment, the invention provides a method of determining the pH of a region of a cell by introducing into the cell a polynucleotide encoding a 20 polypeptide including an indicator having a first fluorescent protein moiety whose emission intensity changes as pH varies between 5 and 10, culturing the cell under conditions that permit expression of the polynucleotide; exciting the indicator; and determining the intensity of the light emitted by the first protein moiety at a first wavelength. The emission intensity of the first fluorescent protein moiety indicates the pH of the region of the cell in which the indicator is present.

The polynucleotide can be introduced using methods described above. Thus, the method can be used to measure intracellular pH in cells cultured in vitro, e.g., HeLa cells, or alternatively in vivo, e.g., in cells of an animal

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carrying a transgene encoding a pH-dependent fluorescent indicator protein.

Fluorescence in the sample can be measured using a fluorometer. In general, excitation radiation, from an 5 excitation source having a first wavelength, passes through excitation optics. The excitation optics cause the excitation radiation to excite the sample. In response, fluorescent proteins in the sample emit radiation which has a wavelength that is different from the excitation wavelength. Collection optics then collect the emission 10 from the sample. The device can include a temperature controller to maintain the sample at a specific temperature while it is being scanned. If desired, a multi-axis translation stage can be used to move a microtiter plate holding a plurality of samples in order to position 15 different wells to be exposed. The multi-axis translation stage, temperature controller, auto-focusing feature, and electronics associated with imaging and data collection can be managed by an appropriately programmed digital computer. The computer also can transform the data collected during 20 the assay into another format for presentation.

Methods of performing assays on fluorescent materials are well known in the art and are described in, e.g., Lakowicz, J.R., Principles of Fluorescence

25 Spectroscopy, New York:Plenum Press (1983); Herman, B., Resonance energy transfer microscopy, in: Fluorescence Microscopy of Living Cells in Culture, Part B, Methods in Cell Biology, vol. 30, ed. Taylor, D.L. & Wang, Y.-L., San Diego: Academic Press (1989), pp. 219-243; Turro, N.J., Modern Molecular Photochemistry, Menlo Park:

Benjamin/Cummings Publishing Col, Inc. (1978), pp. 296-361.

The pH can be analyzed on cells in vivo, or from samples derived from cells transfected with polynucleotides

or proteins expressing the pH indicator proteins. Because fluorescent pH indicator proteins can be expressed recombinantly inside a cell, the pH in an intracellular region, e.g., an organelle, or an extracellular region of an organism can be determined simply by determining changes in fluorescence.

Fluorescent protein pH sensors may vary in their respective pK, and the differences in pK, can be used to select the most suitable fluorescent protein sensor most suitable for a particular application. In general, a sensor protein should be used whose pK, is close to the pH of the sample to be measured. Preferably the pKa is within 1.5 pH unit of the sample. More preferably the pK is within 1 pH unit, and still more preferably the pK, is within 0.5 pH 15 unit of the sample.

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Thus, a fluorescent protein pH sensor having a pKa of about 7.1, e.g., the EYFP mutant described below, is preferred for determining the pH of cytosolic, Golgi, and mitochondrial matrix pH areas of a cell. The YFP-H148G, YFP-H148O, EYFP-H148G and EYFP-H148O mutants are well-suited for measuring the pH of alkaline environments, e.g., mitochondrial matrix, as they have a pKa of 7.5 and 8.0, respectively.

For more acidic organelles, a fluorescence sensor protein having a lower pK, e.g., a pK of about 6.1, is 25 preferred.

To minimize artefactually low fluorescence measurements that occur due to cell movement or focusing, the fluorescence of a fluorescent protein pH sensor can be compared to the fluorescence of a second sensor, e.g., a second fluorescent protein pH sensor, that is also present in the measured sample. The second fluorescent protein pH sensor should have an emission spectra distinct from the

first fluorescent protein pH sensor so that the emission spectra of the two sensors can be distinguished. Because experimental conditions such as focusing and cell movement will affect fluorescence of the second sensor as well as the first sensor, comparing the relative fluorescence of the two sensors allows for the normalization of fluorescence.

A convenient method of comparing the samples is to compute the ratio of the fluorescence of the first fluorescent protein pH sensor to that of the second fluorescent protein pH sensor.

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KITS The materials and components described for use in the methods of the invention are ideally suited for the preparation of a kit. Such a kit may comprise a carrier 15 means being compartmentalized to receive one or more container means such as vials, tubes, and the like, each of the container means comprising one of the separate elements to be used in the method. For example, one of the container means may comprise a polynucleotide encoding a fluorescent 20 protein pH sensor. A second container may further comprise fluorescent protein pH sensor. The constituents may be present in liquid or lyophilized form, as desired. Unless otherwise defined, all technical and scientific terms used herein have the same meaning as 25 commonly understood by one of ordinary skill in the art to which this invention belongs. Although methods and materials similar or equivalent to those described herein can be used in the practice or testing of the present invention, suitable methods and materials are described

30 invention, suitable methods and materials are described below. All publications, patent applications, patents, and other references mentioned herein are incorporated by reference in their entirety. In case of conflict, the

present specification, including definitions, will control. In addition, the materials, methods, and examples are illustrative only and not intended to limit the scope of the invention described in the claims.

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Examples

EXAMPLE 1

CONSTRUCTION OF FLUORESCENT PROTEIN PH SENSORS

Fluorescent protein pH sensors were constructed by engineering site-specific mutations in polynucleotides encoding forms of the Aequora victoria green fluorescent protein (GFP). The starting GFP variant was the polynucleotide encoding the GFP variant EGFP (for enhanced green fluorescent protein). The EGFP variant had the amino acid substitutions F64L/S65T/H231L relative to the wild-type Aequora victoria GFP sequence.

The ECFP (enhanced cyan fluorescent protein) mutant was constructed by altering the EGFP polynucleotide sequence so that it encoded a protein having the amino acid substitutions K26R/F64L/S65T/Y66W/N146I/M153T/V163A/N164H/H231L relative to the wild-type GFP amino acid sequence. A second variant, named EYFP (enhanced yellow fluorescent protein) was constructed by altering the EGFP polynucleotide to encode a protein having the amino acid substitutions S65G/S72A/T203Y/H231L relative to the amino acid sequence of GFP. A third variant, named EYFP-V68L-Q69K, was constructed by altering the EGFP polynucleotide to encode a protein

having the amino acid substitutions S65G/V68L/Q69K/S72A/T203Y relative to the amino acid sequence of GFP.

A HindIII site and Kozak consensus sequence (GCCACCATG) was introduced at the 5' end of the polynucleotide encoding the GFP variants, and an EcoR1 site was added at the 3' end of the gene of each indicator, and the fragments were ultimately ligated into the HindIII/EcoR1 sites of the mammalian expression vector pcDNA3 (Invitrogen). EGFP and EYFP mutant proteins with no targeting signals were used as indicators of pH in the cytosol or nucleus.

To construct fluorescent protein pH sensors to use as pH indicators in the Golgi, polynucleotides encoding the 81 N-terminal amino acids of the type II membrane-anchored protein galactosyltransferase (GT:UDP-galactose- β ,1,4-galactosyltransferase. EC 2.4.1.22) ligated to

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polynucleotides encoding EGFP, ECFP, or EYFP. The polynucleotides encoding the resulting proteins were named GT-EGFP, GT-ECFP, and GT-EYFP, respectively.

Mitochondrial matrix fluorescent protein pH sensors were constructed by attaching polynucleotides encoding 12 amino acids at the amino terminus of the presequence of subunit IV of cytochrome c oxidase (Hurt et al, EMBO J. 4:2061-68 (1985) to a polynucleotide encoding the amino acid sequence Arg-Sea-Gly-Ile, which in turn was ligated to polynucleotides encoding ECFP or EYFP. These constructs were labeled ECFP-mito or EYFP-mito.

The constructs used to examine intracellular pH are summarized in FIG. 1.

EXAMPLE 2

PH TITRATION OF FLUORESCENT SENSOR PROTEINS IN VITRO

The pH sensitivity of the fluorescence of the proteins ECFP, EGFP, EYFP, GT-EGFP, and GT-EYFP was first examined.

Absorbance spectra were obtained in a Cary 3E spectrophotometer (Varian). For pH titration, a monochromator-equipped fluorometer (Spex Industries, NJ) and a 96-well microplate fluorometer (Cambridge Technology) were used. In the latter case the filters used for excitation were 482 ± 10 (460 ± 18 for ECFP) and for emission were 532 ±14. Filters were named as the center wavelength ± the half-bandwidth, both in nm. The solutions for cuvette titration contained 125 mM KCl, 20 mM NaCl, 0.5 mM CaCl₂, 0.5 mM MgCl₂, and 25 mM of one of the following buffers-acetate, Mes, Mops, Hepes, bicine, and Tris.

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EYFP showed an acidification-dependent decrease in the absorbance peak at 514 nm and a concomitant increase in absorbance at 390 nm (FIG. 2A). The fluorescence emission (527-nm peak) and excitation spectra decreased with decreasing pH, but the fluorescence excitation spectrum showed no compensating increase at 390 nm. Therefore, the species absorbing at 390 nm was nonfluorescent. The apparent pKa (pK'a) of EYFP was 7.1 with a Hill coefficient (n) of 1.1 (FIG. 2B).

 $\mbox{EGFP fluorescence also was quenched with decreasing pH. The pK'a of EGFP was 6.15, and n was 0.7. } \\$

The change in fluorescence of ECFP (Tyr66→Trp in the chromophore) with pH was smaller than that of EGFP or EYFP (pK'a 6.4, n, 0.6) (FIG. 2B). The fluorescence change was

reversible in the pH range 5-8.5 for all three proteins, which covers the pH range of most subcellular compartments. These results demonstrate that the GFP variants EGFP, EYFP, and ECFP can be used as fluorescent protein pH sensors.

EXAMPLE 3

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MEASUREMENTS OF pH IN THE CYTOSOL AND NUCLEUS USING FLUORESCENT PROTEIN pH SENSORS

HeLa cells and AT-20 cells grown on glass coverslips were transiently lipo-transfected (Lipofectin M , GIBCO) with polynucleotide constructs encoding EYFP.

Cells were imaged between 2 and 4 days after transfection at 22°C with a cooled charge-coupled device camera (Photometrices, Tucson, AZ) as described in Miyawaki et al., Nature 388:882, (1997). The interference filters (Omega Optical and Chroma Technology, Brattleboro, VT) used for excitation and emission were 440±10 and 480±15 for ECFP; 480±15 and 535±22.5 for EGFP or EYFP. The dichroic mirrors were 455 DCLP for ECFP and 505 DCLP for EGFP or EYFP. Regions of interest were selected manually, and pixel intensities were spatially averaged after background subtraction. A binning of 2 was used to improve signal/noise and minimize photodamage and photoisomerization of EYFP. High KCl buffer plus 5µM each of the ionophores nigericin (Fluka) and monensin (Calbiochem) was used for in situ titrations in living cells. Cells were loaded with cytosolic pH indicators by incubation with 3 µM carboxy-

Fluorescence of HeLa cells transfected with the gene 30 encoding EYFP was diffusely distributed in the cytosol and

washed for 30 minutes, all at 22°C.

SNARF/AM or BCECF/AM (Molecular Probes) for 45 minutes, then

nucleus. This was expected for a protein of the size of GFP $\hat{}$ (27 kDa), which is small enough to pass through nuclear pores.

The fluorescence observed with EYFP was reversible. Perfusion with NH,Cl caused an increase in fluorescence (rise in pH), which reversed upon washing out the NH,Cl. Conversely, perfusion of lactate, which lowers pH, induced a decrease in fluorescence. The decrease in fluorescence was also reversible on wash-out.

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Calibration of fluorescence intensity with pH in situ was accomplished with a mix of the alkali cation/H+ ionophores nigericin and monensin in bath solutions of defined pH and high K+. Fluorescence equilibrated within 1-4 minutes after each exchange of solution. These results demonstrate that EYFP, when present intracellularly, can report pH in the physiological range.

EXAMPLE 4

MEASUREMENT OF pH IN THE MITOCHONDRIAL MATRIX

USING FLUORESCENT PROTEIN pH SENSORS

To measure pH in the mitochondrial matrix using mutant GFP sensor proteins, HeLa cells and neonatal rat cardiomyocytes were transfected with the fluorescent protein pH sensor EYFP-mito. A Bio-Rad MRC-1000 confocal microscope was used for analysis of the targeted protein. Microscopy analysis revealed that the transfected cells showed a fluorescence pattern indistinguishable from that of the conventional mitochondrial dye rhodamine 123.

In situ pH titration was performed with

nigericin/monensin as described in Example 3. Subsequent
addition of the protonophore carbonylcyanide m-

PCT/HS99/12850 WO 99/64592

chlorophenylhydrazone (CCCP) did not change the fluorescence intensity of the cells. This demonstrates that the nigericin/monensin treatment effectively collapsed the pH gradient (ADH) in the mitochondria.

The estimated pHm was 7.98 + 0.07 in HeLa cells (n = 17 cells, from six experiments). Similar pH values were obtained in a HeLa cell line stably expressing EYFP-mito. Resting pH did not change by superfusion of cells with medium 10 mM glucose, which would provide cells with an 10 oxidizable substrate, but 10 mM lactate plus 1 mM pyruvate caused an acidification, which reversed on washout. This can be accounted for by diffusion of protonated acid or by cotransport of pyruvate /H* through the inner mitochondrial membrane. The protonophore CCCP rapidly induced an acidification of mitochondria to about pH 7.

EXAMPLE 5

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MEASUREMENT OF pH IN THE GOLGI LUMEN USING FLUORESCENT PROTEIN DH SENSORS

The type II membrane-anchored protein 20 galactosyltransferase (GT:UDP-galactose-β,1,4galactosyltransferase. EC 2.4.1.22) has been used as a marker of the trans cisternae of the Golgi apparatus (Roth et al., J. Cell Biol, 93:223-29, (1982)). Accordingly, polynucleotide constructs encoding portions of the GT protein fused to the mutant GFP proteins were constructed as described in Example 1 in order to use the GT sequence to target the fluorescent protein pH sensor to the endoplasmic reticulum.

The pH of the Golgi lumen was measured by transfecting HeLa or AT-20 cells with the constructs GT-

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ECFP, GT-EGFP, or GT-EYFP. Bright juxtanuclear fluorescence was observed, with little increase in diffuse staining above autofluorescnce in most cells.

The fluorescence pattern was examined further in 5 double-labeling experiments using rabbit polyclonal αmannosidase II (α-manII) antibody. Double labeling fluorescence was performed as described by McCaffery et al., Methods Enzymol. 257:259-279 (1995). The α-manII antibody was prepared as described in Velasco et al., J. Cell Biol. 122:39-51 (1993). In the double-staining experiments, it was observed that labeling of the medial trans-Golgi marker α-manII overlapped with GT-EYFP fluorescence.

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α-manII was also fused with ECFP, and the pattern of fluorescence obtained upon transfection of the gene was indistinguishable from that of GT-EYFP by light microcopy.

To identify the subcellular localization of GT-EYFP at higher resolution, immunogold electron microscopy was performed on ultra-thin cryosections by using antibodies against GFP. Immunogold labeling of ultra-thin sections was performed as described by McCaffery et al., supra, using 20 rabbit polyclonal anti-GFP antibody or a monoclonal anti-TGN38 antibody.

In double-labeling experiments, GT-EYFP was found in the medial and trans Golgi, although endogenous GT is present in trans Golgi membranes. The difference in 25 localization may occur as a result of overexpression of the GT-EYFP protein.

When protein TGN38 was used as a trans-Golgi network (TGN) marker, its immunogold localization pattern was found to overlap with that of GT-EYFP in the medial/trans-Golgi membranes. The localization data demonstrate that GT-EYFP labels the medial/trans Golqi. Thus, GT-EYFP can be used to identify the pH of this organelle.

The pH titration of GT-EYFP fluorescence in the Golgi region of the cells after treatment with nigericin/monensin was in good agreement with that of EYFP in vitro (see Example 2). Resting pH in HeLa cells was on average 6.58 (range 6.4-6.81, n = 30 cells, 9 experiments). These results also demonstrate that neither fusion with GT nor the composition of the Golgi lumen affects the pH sensitivity of EYFP. Thus, Golgi-targeted EYFP can be used as a local pH indicator.

The effect of various treatments on the pH of the Golgi was next examined using Golgi-targeted EYFP.

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The pH gradient across the Golgi membrane is maintained by the electrogenic ATP-dependent H' pump (V-ATPase). The V-ATPase generates a apH (acidic inside) and $\Delta\psi$ (positive inside), which opposes further H' transport. The movement of counter-ions, Cl' in (or K' out), with H' uptake would shunt the $\Delta\psi$, allowing a larger Δ PH to be generated. These mechanisms were investigated in intact single HeLa cells transfected with GT-EYFP.

The macrolide antibiotic bafilomycin Al has been 20 shown to be a potent inhibitor of vacuolar type H ATPases (V type). In Hela cells expressing GT-EYFP, bafilomycin Al (0.2 µM) increased pH2 by about 0.6 units, to pH 7.16 (range 7.02-7.37, n = 12 cells. This suggests that the H⁺ pump 25 compensates for a positive H+ efflux or leak. The initial rate of Golgi alkalinization by bafilomycin A1 was 0.52 pH units per minute (range 0.3-0.77, n =12 cells), faster than that reported for other acidic compartments such as macrophage phagosomes (0.09 pH/min). Similar results regarding resting pHG and alkalinization by bafilomycin A1 30 were obtained when HeLa cells were transfected with GT-EGFP. Calibration of GT-EGFP in situ also mirrored its in vitro

titration (FIG. 1B). Thus, both EGFP and EYFP are suitable Goldi pH indicators.

EXAMPLE 6

MEASURING INTRACELLULAR PH WITH TWO

FLUORESCENT PROTEIN SENSORS

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Quantitative measurements of fluorescence with nonratiometric indicators can suffer from artifacts as a result of cell movement or focusing. To correct for these effects, the cyan-emitting mutant GT-ECFP was co-transfected into cells along with GT-EYFP. ECFP has excitation and emission peaks that can be separated from those of EYFP by appropriate filters. In addition, ECFP is less pH-sensitive than EYFP (see FIG. 2B).

FIG. 3A demonstrates that the fluorescence of ECFP changed less than that of EYFP during the course of the experiment. Although the ratio of EYFP to ECFP emission varied between cells, probably reflecting a different concentration of GT-EYFP and GT-ECFP in the Golgi lumen, it changed with pH as expected (FIG. 3B). Bafilomcin Al raised the GT-EYFP/GT-ECFP emission ratio, i.e, it raised pH₀.

EXAMPLE 7

CONSTRUCTION OF YFP THR H148G AND YFP H1480 MUTANTS

The YFP H148G mutant was prepared using as a template a nucleic acid encoding the YFP mutation 10c, which includes the mutations S65G/V68L/S72A/Q80R/T203Y and is described in Ormō et al., Science 273:1392-95 (1997). The YFP H148G mutant was constructed using the PCR-based QUIKCHANGE™ Site-Directed Mutagenesis Kit (Stratagene, La Jolla, CA) following the manufacturer's instructions. The

YFP H148Q mutation was similarly constructed from a nucleic acid encoding the 10C mutation.

The pKa of the YFP H148G mutant was found to be 8.0, while the YFP H148O mutant was found to have a pKa of 7.5.

EXAMPLE 8

EXPRESSION OF mito-YFP H148G IN THE MITOCHONDRIAL MATRIX AT A ph RANGE OF 7.0 TO 8.4

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The high pKa of the mutant YFP H148G allows it the to be used for the precise measurement of mitochondrial matrix pH both in cells at rest and in cells subject to manipulations that decrease mitochondrial pH.

This was demonstrated directly by transfecting a nucleic acid encoding mito-YFP H148G into HeLa cells using the procedures described in Example 4. YFP H148G expression was monitored by observing fluorescence over time.

Mitochondrial pH was also monitored by pH tirration as described in Example 3 using nigericin and monensin.

FIG. 4 shows that HeLa cells transfected with YFP
10 H148G in the mitochondrial matrix were fluorescent at an
initial pH of 8.0 to 8.1 (where measurements began at t ≈ 0
seconds). 5μM CCCp was added at about t ≈ 300 seconds.
Although addition of 5μm CCCP rapidly lowered the pH to 7.0,
fluorescence of mito-YFP H148G was still detectable. Then a
25 calibration was performed by perfusing the cells with
extracellular medium of ph 7, 7.5, 8, and 8.35 containing
the ionophores nigericin plus momensin to equilibrate
mitochondrial pH and extracellular pH. Fluorescence in
mitochondria increased stepwise with each change of
30 extracellular pH.

Fluorescence was also examined, and pH measured, in primary cultures of chick skeletal myotubes transfected with

the mito-YFP H148G mutant. FIG. 5 demonstrates that fluorescence was detectable in the mitochondrial matrix of chicken skeletal myotubes, which had a pH of 8.0-8.1 ($t \approx 0$). Fluorescence was still detectable following addition of 25 μ M forskolin, which did not affect the pH, and after addition of 2 μ M CCCP at t=750 seconds, although CCCP caused the pH to rapidly drop to 6.9 at t=1400 seconds. Thereafter fluorescence continued to be observed during calibration at ph 6.9, 7.6 and 8.0.

These results demonstrate mito-YFP H148G fluorescence is detectable in the mitochondrial matrix over the pH range of 7.0 to 8.4 in both established cell lines (HeLa cells) and primary cultures (chick skeletal myotubes).

EXAMPLE 9

15 EXPRESSION OF mito-YFP H148Q IN RESPONSE TO PH CHANGES

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The YFP H148Q mutant has a pKa of about 7.4, which is intermediate between the pKa of EYFP and YFP mutant H148G. To demonstrate that YFP H148Q can also be used to measure mitochondrial matrix pH, a nucleic acid encoding mito-YFP-H148Q was transfected into HeLa cells. Fluorescence was measured over time (beginning at t \sim 0), including following the addition of 10 $\mu\mathrm{M}$ nigericin in high KCL titration buffer at t \sim 500 seconds.

FIG. 6 reveals the effect of changing mitochondrial pH to 6.9, 7.5, 8, and 8.4 with the ionophore nigericin on fluorescence intensity. Fluorescence decreased to about 175 units at t=1000 seconds by addition of nigericin, which lowered the pH to about 6.9. Fluorescence then returned stepwise to 400 units with each change of the extracellular

medium. These results demonstrate that the fluorescence of

the mito-YFP H148Q mutant can be used to measure the pH of

EXAMPLE 10

MEASURING INTRACELLULAR DH BY COEXPRESSION OF YFP H148G AND 5 A SECOND PH-INSENSITIVE SENSOR

As is discussed above in Example 6, for quantitative measurements it is desirable to compute the fluorescence of the sensor used to measure pH with the fluorescence of a second sensor molecule whose fluorescence does not change 10 over the pH range being tested. A ratiometric measurement is useful to correct for movement or focusing artifacts that may occur during live cell imaging experiments.

To identify a GFP sensor protein suitable for use as a reference protein for measuring mitochondrial matrix pH, the GFP must T203I was expressed in the mitochondria of HeLa cells. The GFP T203I mutant can be excited with light of 400 nm, which does not appreciably excite the pH sensitive YFP mutants.

Fluorescence of HeLa cells transfected with the GFP
20 T203I mutant was monitored for about 400 seconds using an
excitation ratio of 480 nm/400 nm. 10 µM CCCP was then
added to the cells, and fluorescence was monitored for an
additional 250 seconds. Addition of CCCP did not affect
fluorescence. In control experiments, it was observed that
25 addition of CCCP corresponded to a drop in pH of about 1
unit. Thus, the GFP T203I mutant is suitable for use as a
reference, pH-insensitive mutant.

HeLa cells were then transfected with the GFP T203I mutant and YFP H148G. FIG. 7 shows the change of

30 mitochondrial pH with oligomycin and the uncoupler CCCP as the ratio of YFP H148G emission and GFP T203I emission, with excitation of 490 and 400 nm, respectively.

EXAMPLE 11

STRUCTURAL CHARACTERIZATION OF YFP T203Y/S65G/V68L/S72A/H148G

The green fluorescent protein (GFP) from the

5 jellyfish Aequorea victoria has been used extensively in
molecular biology as a fluorescent label. The structures of
WT GFP (Yang et al., Nature Biotech. 14:1246-51, 1996; Brejc
et al., Proc. Natl. Acad. Sci. USA. 94:2306-11, 1997) and
the variant S65T were determined in 1996 (Ormō et al.,

10 Science 273:1392-95, 1996).

A large number of mutants have been identified that exhibit broadly varying absorption and emission maxima (Heim et al. above;, Heim et al., Curr. Biol. 6: 178-82, 1996). The yellow fluorescent protein (YFF) mutant is of particular interest since its spectrum is shifted enough to render it

- interest since its spectrum is shifted enough to render it readily distinguishable from the spectrum of Cyan Fluorescent Protein (CFP) for FRET measurements (Tsien, Ann. Rev. Biochem. 67:509, 1998; Miawaki et al., Nature 388:882-87, 1997). WT GFP exhibits two absorption maxima, where
- 20 the major band absorbs at 398 nm and the minor band at 475 nm (Morise et al., Biochemistry 13:2656-62, 1974). Excitation of either of these bands leads to emission of green light with a maximum between 504 and 508 nm (FIG. 8). Before a structure was available, GFP variants with altered
- 25 spectral characteristics were identified by random mutagenesis. Some of these mutants, such as Y66H and Y66W (Tsien, Ann. Rev. Biochem. 67:509, 1998, Heim et al., Proc. Natl. Acad. Sci. (USA) 91:12501-04) result in blue-shifted absorbance and emission maxima. Others focus on changes in the immediate environment of the chromophore π system, such as S65T (Heim et al., Nature 373: 663-64, 1995) and T2031
 - (Heim et al., Proc. Natl. Acad. Sci. (USA) 91:12501-04, 1994). At physiological pH, S65T exhibits only one major

absorption band at 489 nm, red-shifted by 14 nm from WT GFP, and is almost six times brighter (Heim et al., Nature 373: 663-64, 1995). Yet, the emission spectrum is shifted by only 3 nm to 511 nm, and so cannot easily be distinguished from the wild-type emission. Random mutagenesis techniques produced only one further red-shifted variant, S65T/M153A/K238E, which increases the excitation and emission wavelengths of S65T by 15 and 3 nm respectively (Heim et al., Curr. Biol. 6: 178-82, 1996). Here is described crystal structures of the first set of GFP variants rationally designed based on the x-ray structure of GFP S65T (Ormö et al., Science 273:1392-95, 1996). These variants, termed YFPs (Yellow Fluorescent Proteins), exhibit the longest wavelength emissions of all GFPs generated by mutagenesis (FIG. 8). The YFPs fluoresce around 528 nm, red-shifted by 16 nm as compared to S65T and are easily distinguishable from S65T on a fluorescence microscope.

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The specific YFP investigated is the quadruple-mutant T203Y/S65G/V68L/S72A, where the substitution T203Y was introduced based on the structural 20 considerations detailed below and is believed responsible for the red-shift. The other three mutations have been shown to improve its brightness in live cells (Cormack et al., Gene 173:33, 1996). The T203Y mutation would have been difficult to identify by random mutagenesis since this amino 25 acid substitution requires three substitutions at the nucleotide level. Since Thr203 is positioned close to the chromophore, it was postulated that its replacement with a tyrosine would result in π -stacking interactions between the 3.0 chromophore and the highly polarizable phenol (Ormö et al., Science 273:1392-95, 1996), leading to red-shifted spectral properties. The structure of S65T suggested that an aromatic amino acid introduced in position 203 would extend

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into the water-filled cavity adjacent to the chromophore (Ormö et al., Science 273:1392-95, 1996). Replacement of Thr203 with any of the aromatic amino acids His, Trp, Tyr, or Phe was found to lead to the desired spectral shifts (Ormo et al., Science 273:1392-95, 1996, Dickson et al., Nature 388:355-58, 1997). The most dramatic red-shift was observed for the T203Y substitution, therefore this variant has been termed YFP.

In order to determine the role of His148 in modulating the pKa of the chromophore or its spectral 10 properties, an additional mutation, H148G, was introduced into the YFP background. The x-ray structures of YFP and YFP H148G were analyzed in order to better correlate structural changes with spectral properties. GFP variants were prepared as described in Example 6, above. This 15 template incorporates the mutations T203Y/S65G/V68L/S72A, as well as the ubiquitous Q80R substitution that was accidentally introduced into the gfp cDNA early on (Ormö et al., Science 273:1392-95, 1996; Chalfie et al., Science 263:802-05, 1994). All GFP variants were expressed and 20 purified as described (Ormö et al., Science 273:1392-95,

1996).

Structural Determination of YFP H148G

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YFP H148G was concentrated to 12 mg/ml in 20 mM HEPES pH 7.9. Rod-shaped crystals with approximate dimensions of $1.8 \times 0.08 \times 0.04$ mm were grown in hanging drops containing 2 µl protein and 2 µl mother liquor at 4°C within four days. The mother liquor contained 16% PEG 4000, 50 mM sodium acetate pH 4.6, and 50 mM ammonium acetate.

X-ray diffraction data were collected from a single crystal 3.0 at room temperature using a Xuong-Hamlin area detector (Hamlin, Methods. Enzymol. 114:416-52, 1985). Data were

collected in space group P2,2,2, to 99% completeness at 2.6 Å resolution, and reduced using the supplied software (Howard et al., Methods Enzymol. 114:452-71, 1985). Unit cell parameters were a = 52.0, b = 62.7, and c = 69.9. The GFP S65T coordinate file (Ormo et al., Science 273:1392-95, 1996) which served as a model for phasing was edited to reflect the mutations, with the introduced residues Tyr203 and Leu68 initially modeled as alanines to prevent model bias. A model for the anionic chromophore was obtained by semi-empirical molecular orbital calculations using AM1 in 10 the program SPARTAN version 4.1 (Wavefunction Inc., Irvine, CA). The minimized structure, which was planar, compared very favorably with a related small molecule crystallographic structure (Tinant et al., Cryst. Struct. Comm. 9:671-74, 1980), and also with the model used during 15 refinement of GFP S65T, where a simpler modeling program had been employed (Ormo et al., Science 273:1392-95, 1996). Using the program TNT (Tronrud et al., Acta Crystallogr. Sect. A 43:489, 1987), rigid body refinement was carried out to position the isomorphous model in the 20 unit cell of YFP H148G. Initial positional refinement was carried out using the data to 4.0 Å, then to 3.5, 3.0, and finally to 2.6 Å. Electron density maps (2Fo - Fc and Fo -

carried out using the data to 4.0 Å, then to 3.5, 3.0, and finally to 2.6 Å. Electron density maps (2Fo - Fc and Fo - Fc) were inspected using O (Tronrud et el., above), and solvent molecules were added if consistent with Fo-Fc features, and only when in proximity of hydrogen bond partners. B-factors were refined using a strong correlation between neighboring atoms due to the relatively low resolution. Since no B-factor library is available for the chromophore itself, the B-factors of all chromophore atoms were set to the values obtained in the 1.9 Å structure of GFP S65T (Ormö et al., Science 273:1392-95, 1996), and then

refined as a group, with identical shifts for the grouped atoms.

Structure Determination of YFP

asymmetric unit.

YFP was concentrated to 10 mg/ml in 50 mM HEPES pH 7.5. After 2 weeks crystals grew to a size of 0.03 x 0.12 x 0.8 mm at 15°C in hanging drops containing 5 | 1 protein and 5 | 1 well solution, which contained 2.2 M sodium/potassium phosphate pH 6.9. These crystals belong to space group P2,2,2 and have the unit cell dimensions a = 77.1, b =117.4, w and c = 62.7. X-ray diffraction data were collected on 10 two isomorphic crystals at room temperature using an Raxis-IV imaging plate mounted on a Rigaku RUH3 rotating anode generator equipped with mirrors. The data were processed with Denzo and scaled using ScalePack (Otwinowski et al., Methods Enzymol. 276:307-26, 1997). The YFP 15 structure was solved by molecular replacement using the program AMoRe (Navaza, Acta Crystallogr. A50:157-63, 1994), with the 1.9 A GFP S65T coordinate file as the search model (Ormö et al., Science 273:1392-95, 1996). Two solutions were identified, consistent with two molecules per 20

For refinement, the 2.6 Å structure of YFP H148G was chosen as the initial model, which was edited to reflect the mutations present in YFP. To avoid model bias, the occupancies of the Tyr203 side chain atoms and all chromophore atoms were set to zero during the first several rounds of refinement. Constrained NCS averaging over the A and B chains in the asymmetric unit was applied, initial refinement was carried out to 3.5 Å only, and the electron density maps (2Fo - Fc and Fo -Fc) were averaged. These maps were then inspected, and the model adjusted using 0 (Jones et al., Acta Crystallogr. Sect. A 47:110, 1991), followed by additional positional refinement to 2.5 Å.

Chromophore and Tyr203 densities were very clear, and both were planar. The model was edited to include these groups in refinement, and solvent molecules were added where appropriate. B-factors were refined using a strong correlation between neighboring atoms due to the relatively low resolution.

Comparison of the Structure of YFP and YFP H148G

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YFP crystallized in 2.2 M Na/K phosphate at pH 6.9 in spacegroup P2,2,2, with 2 molecules per asymmetric unit (chains A and B). The GFP S65T structure was used as a search model for molecular replacement against a 3.0 Å dataset using the program AMoRe (Navaza, Acta Crystallogr. A50:157-63, 1994), and the structure was refined. Later, the refined structure of YFP H148G (see below) was used for phasing and refinement against a 2.5 Adataset. Even though the introduced Tyr203 and the chromophore itself were not modeled during early cycles of refinement, clear electron density for a planar chromophore and a stacked Tyr203 phenol was immediately apparent. Non-crystallographic symmetry (NCS) constraints were employed throughout refinement of the model using TNT, and maps were averaged. At the end of refinement, non-averaged maps for the A-and B-chain in the asymmetric unit were calculated and compared to each other. No obvious features were identifiable that would suggest significant differences between the two chains. A test run of refinement without any NCS constraints confirmed that the differences would be smaller than the rms error of a 2.5 Å structure. Therefore, the NCS constraints were not relaxed or eliminated. Data collection and atomic model statistics are shown in Table 33. The final R-factor of the YFP model

was 19.2% for all data between 20 and 2.5 Å resolution.

Table 33. Data collection and atomic model statistics of YFP and YFP H148G.

		YFP	YFP H148G
10	Total observations Unique reflections Completeness* Completeness (shellb) Number of crystals *merge* (%)	53,039 18,916 92% 94% 2 8.0%	29,904 7,373 99% 97% 1 6.5%
	Resolution Atomic model statistics: Spacegroup F2.2.2,	2.5 Å P2 ₁ 2 ₁ 2	2.6 Å
15	Molecules per asymm, unit Crystallographic R-factor Protein atoms Solvent atoms per asymmm, unit Bond length deviations (Å)	2 0.192 1,810 130 0.013	1 0.159 1,810 30 0.012
20	Bond angle deviations (°) Thermal parameter restraints (Å ²)	1.76 4.53	2.07 3.82

 $^{^{\}circ}$ Completeness is the ratio of the number of observed I > 0 divided by the theoretically possible number of intensities.

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The refined YFP structure clearly shows that the overall fold is undisturbed, with an rms deviation from the GFP S65T structure of 0.36Å for α -carbons. Three larger contact areas with adjacent molecules were identified. The largest of these covers about 722 Å2 of one monomer surface, includes a series of hydrophobic residues consisting of

5 Ala206, Phe223, and Leu221, and also a number of hydrophilic contacts. This interface is essentially identical to the

 $^{^{\}rm b}$ Shell is the highest resolution shell (2.56 to 2.50 Å for YFP, and 2.80 to 2.60 Å for YFP H148G)

 $^{^{\}rm c}$ R_{\rm merge} = Σ | I_{hk1} -<1>| / Σ < I > where < I > = average of individual measurements of $I_{\rm hk1}$.

dimer interface for WT GFP described by Yang et al., Nature Biotech. 14:1246-51, 1996. High salt conditions during crystallization experiments appear to favor dimerization, as has been suggested previously (Palm et al., Nat. Struct. Biol. 4:361-65, 1997).

YFP H148G crystallized as a monomer in the presence of polyethylene glycol and acetate at pH 4.6 in spacegroup P2.2.2. isomorphous to S65T (Ormö et al., Science 273:1392-95, 1996) and the blue-emission variant BFP (Wachter et al., Biochemistry 36:9759-65, 1997). Molecular replacement using 10 the S65T structure for phasing and refinement gave a final model with an R-factor of 15.9% for all data between 24.0 and 2.6 Å (Table 33). As with YFP, electron density for a stacked phenol was clearly visible even before the Tyr203 ring was modeled. The rms deviation between YFP H148G and S65T a-carbons is 0.31 Å, and the deviation between YFP H148G and YFP a-carbons is 0.35 Å. The b-strands of the two YFP variants overlay closely in all areas except around the C. of residue 148 where a movement of 1.1 Å is observed. This movement has not been observed in other pH 4.6 structures grown under similar conditions and crystallizing in the same space group, such as the BFP structure (Wachter et al., Biochemistry 36:9759-65, 1997). Residue 148 and

adjacent residues are not involved in crystal contacts, further indicating that the observed movement is due to the H148G substitution, not crystallization conditions.

π -Stacking of the introduced phenol

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Electron densities of the chromophore and the phenol ring of Tyr203 appeared to be completely planar before the atoms for these groups were added to the model. When the tyrosine side chain was first introduced into the model, it was modelled as co-planar to the chromophore. Refinement

consistently rotated the phenol ring by 12 with respect to the chromophore plane in both YFP and YFP H148G. FIG. 9 shows the electron density of the refined YFP chromophore structure together with the phenol ring of Tyr203. The distance of the closest approach between atoms of the two interacting rings is 3.3 to 3.4 Å, and occurs at that edge of the chromophore plane that is opposite the exo-methylene bond (FIG. 9). It appears that the phenol tilts towards this area of the chromophore since it is more open, with fewer atoms to clash with sterically.

The distance of largest separation between the rings is 3.5 to 3.8 Å, and occurs at the opposite edge, where steric clash with the exo-methylene carbon could occur. This range of plane-to-plane distances is typical for

15 face-to-face π to π stacking interactions found in proteins, and consistent with interaction energy calculations that show a potential energy minimum for two horizontally stacked benzene molecules with a vertical separation of 3.3 Å (Burley et al., J. Am.Chem. Soc. 108,

7995-8001, 1986). A recent analysis of protein structures has led to the conclusion that aromatic ring interactions in an off-centered parallel orientation have an energetically favorable, stabilizing effect, and in fact are the preferred interactions (McGaughey et al., J. Biol. Chem. 273,

25 15458-63, 1998).

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Positional shift of the chromophore

The entire chromophore ring system of YFP has moved out towards the protein surface by about 0.9 Å when compared to S65T or WT GFP. The chromophore of YFP H148G has moved in the same direction but to a lesser extent, about 0.5 Å. Overlay of all $\alpha\text{-carbons}$ shows that this shift is very much a local effect, only involving residues 65 to 68. The

overlay suggests that this shift may be due to the compensating effects of the V68L and S65G substitutions. The Leu68 Cδ1 occupies the same space as the original Val68 Cyl. whereas the Leu68 backbone is displaced so that the chromophore is pushed further out towards the protein surface. As part of the same movement, the $C\alpha$ of Gly65 is pushed into the position of the wild-type C_g of Ser65. The V68L and S65G substitutions had been previously found to significantly increase the brightness of GFP-expressing cells (Cormack et al., Gene 173:33, 1996) in a WT background, and were suggested to either improve folding at 37° or increase the rate of chromophore formation. It is unclear at this point why the chromophore is not shifted to the same extent in the YFP and YFP H148G structures, though both of them incorporate the V68L and S65G mutations. Even though the imidazolinone ring of the YFPs is not in the same position as in WT GFP (Brejc et al., Proc. Natl. Acad. Sci. USA. 94:2306-11, 1997) , S65T (Ormö et al. Science 273:1392-95, 1996), and blue-fluorescent protein BFP

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Acad. Sci. USA. 94:2306-11, 1997), S65T (Ormo et al. Science 273:1392-95, 1996), and blue-fluorescent protein BFP (Wachter et al., Biochemistry 36:9759-65, 1997), no electron density consistent with partially formed or unformed chromophore is observed. This indicates that the machinery to generate the chromophore is not only intact, but more flexible than previously thought. Apparently, the exact positions of the backbone atoms of residues 65 and 67 that undergo the cyclization reaction is not as crucial as was previously suggested, based on the nearly exact superposition of the imidazolinone rings observed in WT GFP, S65T, and BFP (Yang et al., Nature Biotech. 14:1246-51, 1996) Protein K. et al. Proc. Natl. Acad. Sci. USA 94:2306-

30 1996; Brejc, K. et al., Proc. Natl. Acad. Sci. USA 94:2306-2311, 1997; Ormö et al., Science 273:1392-95, 1996; Palm et al., Nat. Struct. Biol. 4:361-65, 1997; Wachter et al., Biochemistry 36:9759-65, 1997).

Chromophore spectral properties, charge state and hydrogen bonding interactions $% \left(1\right) =\left(1\right) +\left(1\right) +\left($

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The spectral properties of the YFPs were examined. Small aliquots of protein (16 mg/ml) were diluted 48-fold into 75 mM buffer (acetate, phosphate, Tris, or CHES), 140 mM NaCl, and then scanned for absorbance between 250 and 600 nm (Shimadzu 2101 spectrophotometer at medium scan rate and room temperature). The optical density at 514 or 512 nm was plotted as a function of pH and computer-fitted to a titration curve (Kaleidagraph*, Synergysoftware).

Fluorescence measurements were carried out on a Hitachi F4500 fluorescence spectrophotometer at a constant protein concentration of approximately 0.01 mg/ml, with buffer conditions identical to those of absorbance measurements. The excitation wavelength was set to the absorbance maximum of the long-wave band of the particular mutant. The emission was scanned between 500 and 600 nm, and peak emission intensity was plotted as a function of pH and curve-fitted.

Like S65T (Kneen et al., Biophys. J. 74:1591-99, 1998), the YFPs have two absorbance maxima whose relative ratio is pH-dependent (FIG. 8 and Table 34). The UV absorption peaks at 392 (YFF) or 397 nm (YFF H148G) have been ascribed to the neutral chromophore, whereas the visible absorption peaks at 514 (YFF) or 512 nm (YFP H148G) have been ascribed to the anionic chromophore (Niwa et al., Proc. Natl. Acad. Sci. (USA) 93: 13617-22, 1996). The lower energy peak exhibits clear vibrational structure as indicated by the pronounced shoulder at 480-490 nm, and its mirror-image relationship with the emission band is striking (FIG. 8). These features are consistent with luminescence properties of large and rigid systems in condensed phases

(Barltrop et al., Principles of Photochemistry, John Wiley and Sons, New York, 1978, pp. 51-52 and 78-79), and may be more pronounced in the YFPs due to decreased chromophore flexibility in the presence of the stacked phenol. YFPs fluoresce intensely when excited at the longer-wavelength band, with maximum emission occurring at 528 nm (FIG. 8). Fluorescence is extremely weak when the excitation occurs at the shorter-wavelength band (Table 34), even if the experiment is carried out at a pH where this peak dominates. The chromophore pKa in the intact protein 10 was determined to be $7.00(\pm0.03)$ for YFP and $8.02(\pm0.01)$ for YFP H148G by absorbance measurements at varying pH. The pKa values determined by fluorescence were 6.95 (±0.03) and 7.93 (±0.04), respectively, for the two variants. The YFP 15 pKa is remarkably similar to that of

Table 34. Summary of Absorption and Emission Maxima.

	absorbance	absorbance	emission ^a	emission ^a
	band #1	band #2	band	band
WT GFP	398	475	460/508	504
S65T	394	489	(weak)	511
YFP	392	514	(weak)	528
HFP H148G	397	51122	(weak)	528

^a The emission band #1 results from excitation at the absorbance peak #1, and the emission band #2 results from excitation at the absorbance peak #2.

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EYFP(S65G/S72A/T203Y/H231L). All titration curves gave an excellent fit to a single pKa value.

It is likely that the charge state of the chromophore is mixed in the YFP crystals which were grown at pH 7, and which is the chromophore pKa. In YFP, Hisl48 is directly hydrogen-bonded to the phenolic end of the chromophore. Its

electron density is well-defined, suggesting that the imidazole ring does not change position when the chromophore ionizes. It is therefore unlikely that structural rearrangements in the immediate chromophore environment occur in response to changes in chromophore charge state. In both the YFP and YFP H148G structures, the phenolic end of the chromophore is nearly in H-bonding contact with bulk solvent via two ordered waters, and therefore may not be as tightly embedded in the protein as in WT and S65T (Brejc et al., Proc. Natl. Acad. Sci. USA. 94:2306-11, 1997, Ormő et al., Science 273:1392-95, 1996]. Structural readjustments to accommodate the anion may only affect solvent molecules.

The strong hydrogen bond to Arg96 that has been

suggested to play a role in the chemistry of backbone

cyclization (Ormö et al., above) is maintained in both
structures. The carbonyl oxygen of the chromophore
imidazolinone ring interacts with two hydrogen bond donors,
Arg96 and Gln69 in YFP, and Arg96 and Gln94 in YFP H148G.
This compares to similar interactions with Arg96 and Gln94

in WT and S65T. The Glu222 carboxy oxygen approaches the
chromophore imidazolinone ring nitrogen to within 3.0 (YFF)
and 3.3 Å (YFF H148G), considerably closer than in WT and
S65T (4.3 and 4.0 Å, respectively). This close approach

appears to be related to the chromophore positional shift described above. Distance and geometry for hydrogen bonding between Glu222 and the chromophore ring nitrogen are excellent in YFP, and somewhat less optimal in YFP H148G, where the presumed H-bond makes roughly a 45° angle with the chromophore plane. The YFP structure is the first GFP structure solved that suggests H-bonding interactions of the

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structure solved that suggests H-bonding interactions of the heterocyclic ring nitrogen originating from Tyr66. The most likely interpretation in terms of charge states is a deprotonated ring nitrogen and a protonated Glu222,

rendering both groups neutral, however, it is clear that they share a proton.

Solvent-accessible surface and cavities

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The mutation H148G was introduced into YFP to examine the effects of solvent accessibility on the fluorescent properties and the ionization constant of the chromophore. In all GFP structures examined to date, the β -barrel is somewhat perturbed around the phenolic end of the chromophore. The β -strand that covers the chromophore in that area bulges out around His148, so that the backbone from residue 144 to 150 is not directly hydrogen-bonded to the adjacent backbone between residues 165 and 170. Rather, they are laced together by forming H-bonds with the imidazole ring of His148 (Arg168 backbone N to His148 N., in S65T and WT GFP) and several water molecules. The phenolic end of the chromophore is located directly "behind" the ring of His148. It was anticipated that substitution of His with Gly would open up a solvent channel to the chromophore in the absence of other structural perturbances, or perhaps to permit the bulge to close.

The crystal structure clearly shows this anticipated solvent channel as an invagination of the protein surface with no ordered solvent molecules within. Elimination of the imidazole ring in the H148G substitution only leads to minor structural rearrangements of protein groups. The $\beta\text{-strands}$ do not close up to form a directly H-bonded sheet between residues 144 and 150. Instead, the C α of residue 148 has actually moved in the opposite direction by 1.1 Å, causing an even larger strand separation between the

backbones of residues 148 and 168. The side chain of Ile167 has moved by 1.1 Å towards the space previously occupied by the imidazole ring. Nevertheless, direct solvent access to

the phenolic end of the chromophore is greatly improved. Calculation of the solvent-accessible area of the chromophore using a probe sphere of radius of 1.4 A (Connolly, Science 221:709-13, 1983), as implemented by UCSF MidasPlus (UCSF MidasPlus®, Computer Graphics Laboratory, University of San Francisco, CA 94143), shows that 22% of the chromophore surface is solvent-accessible. Only the phenolic end of the chromophore is exposed to exterior solvent, though, due to the opening in the protein wall. The phenolic oxygen of the chromophore is also hydrogen-bonded to a water molecule that is near H-bonding distance to a surface water, though a 1.4 Å probe cannot access the chromophore via this path. If both the solvent channel as well as this hydrogen bond are included, 8% of the chromophore surface is accessible to exterior solvent, entirely at the phenolic end, and 14% is accessible to interior solvent due to contact with internal cavities.

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YFP H148G contains two larger interior cavities that are in contact with the chromophore cavity and filled with some ordered waters. The cavity that was largest in S65T has decreased in size from approximately 127 Å3 (S65T) to 88 Å3 (YFP H148G), because some of the space is now filled with the phenol of Tyr203. In YFP, this cavity is not accessible to a 1.4 Å probe at all since several groups have moved into this space. The more significant structural adjustments are C,2 of Val224, which has moved by 1.4 Å, and C₆₁ of Leu42 which has moved by 2.0 Å, essentially filling the cavity. The second larger cavity in contact with the chromophore is nearly invariant for S65T, YFP, and YFP H148G, and is

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Solvent accessibility to the chromophore

YFP H148G was found to be highly fluorescent, with bright greenish-yellow color under ordinary day light. The light-emitting properties of the fluorophore do not appear 5 to be changed to any extent by the introduction of a solvent channel to the chromophore, indicating that significant quenching does not occur.

Since the protein fold is entirely intact in YFP H148G in spite of the generation of an opening in the β -barrel, the H148G substitution may be especially useful for allowing 10 access of various small-molecule species to the chromophore. This substitution may be introduced into other GFP variants with a larger cavity adjacent to the chromophore, such as S65T [7] or S65G, allowing for analyte binding studies where specific spectral shifts due to the interaction with small 15 molecules or ions of interest could be monitored. The highest ionization constant of all variants examined to date is found for the YFP H148G mutant with a pKa of 8.0. In this mutant, the chromophore is solvent exposed, consistent with a similarly high pKa when the protein is denatured 20

(Nageswara et al., Biophys. J. 32:630-32, 1980).

Other Embodiments

It is to be understood that while the invention has been described in conjunction with the detailed description 25 thereof, the foregoing description is intended to illustrate and not limit the scope of the invention, which is defined by the scope of the appended claims. Other aspects, advantages, and modifications are within the scope of the following claims. 30

What is claimed is:

1 1. A functional engineered fluorescent protein
2 whose amino acid sequence is substantially identical to the
3 amino acid sequence of Aequora green fluorescent protein
4 (SEQ ID NO:__) and whose emission intensity changes as pH
varies between 5 and 10.

- 2. The functional engineered fluorescent protein of claim 1, wherein the amino acid sequence of the protein includes the substitutions S65G/S72A/T203Y/H231L in the amino acid sequence of Aequora green fluorescent protein (SEQ ID NO:).
- 3. The functional engineered fluorescent protein of claim 1, wherein the amino acid sequence of the protein includes the substitutions S65G/V68L/Q69K/S72A/T203Y in the amino acid sequence of Aequora green fluorescent protein (SEC ID NO:__).
- 4. The functional engineered fluorescent protein of claim 1, wherein the amino acid sequence of the protein includes the substitutions

 K26K/F64L/S65T/Y66W/N146I/M153T/V163A/N164H/H231L in the amino acid sequence of Aeg

(SEO ID NO:).

- 5. The functional engineered fluorescent protein of claim 1, wherein the amino acid sequence of the protein includes the substitution H148G in the amino acid sequence of Aeguera green fluorescent protein (SEQ ID NO:).
- 1 6. The functional engineered fluorescent protein of 2 claim 1, wherein the amino acid sequence of the protein

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3 includes the substitution H148Q in the amino acid sequence of Aequora green fluorescent protein (SEQ ID NO:).

- 7. A polynucleotide comprising a nucleotide
- 2 sequence encoding a functional engineered fluorescent
- 3 protein whose amino acid sequence is substantially identical
- to the amino acid sequence of Aequora green fluorescent
- 5 protein (SEQ ID NO:__) and whose emission intensity changes as pH varies between 5 and 10.
- 1 8. The polynucleotide of claim 7, wherein the amino 2 acid sequence of the protein includes the substitutions 3 S65G/S72A/T203Y/H231L in the amino acid sequence of Aequora
- 9. The polynucleotide of claim 7, wherein the amino
 call acid sequence of the protein includes the substitutions
 S65G/V68L/069K/872A/T203Y in the amino acid sequence of
- Aequora green fluorescent protein (SEQ ID NO:__).

green fluorescent protein (SEQ ID NO:).

- 1 10. The polynucleotide of claim 7, wherein the 2 amino acid sequence of the protein includes the
- 3 substitutions
 - K26R/F64L/S65T/Y66W/N146I/M153T/V163A/N164H/H231L in the
- 5 amino acid sequence of Aequora green fluorescent protein (SEQ ID NO:__).
- 1 11. The polynucleotide of claim 7, wherein the
- 2 amino acid sequence of the protein includes the substitution
- 3 H148G in the amino acid sequence of Aequora green fluorescent protein (SEQ ID NO:).

1 12. The polynucleotide of claim 7, wherein the 2 amino acid sequence of the protein includes the substitution 3 H148Q in the amino acid sequence of Aequora green fluorescent protein (SEO ID NO:).

- 1 13. An expression vector comprising expression
 2 control sequences operatively linked to a polynucleotide
 3 molecule comprising a nucleotide sequence encoding a
 4 functional engineered fluorescent protein whose amino acid
 5 sequence is substantially identical to the amino acid
 6 sequence of Aequorea green fluorescent protein (SEQ ID NO:2)
 7 and whose emission intensity changes as pH varies between 5
- 1 14. A recombinant host cell comprising the expression vector of claim 13.
- 1 15. The recombinant host cell of claim 14, wherein the recombinant host cell is a prokaryotic cell.
- 1 16. The recombinant host cell of claim 14, wherein the recombinant host cell is a eukaryotic cell.
- 1 17. A method for determining the pH of a sample comprising:
- 3 contacting the sample with an indicator comprising a
- 4 first fluorescent protein moiety whose emission intensity
- 5 changes as pH varies between pH 5 and 10;
- 6 exciting the indicator; and

and 10.

- 7 determining the intensity of light emitted by the
- 8 first fluorescent protein moiety at a first wavelength,
- 9 wherein the emission intensity of the first fluorescent protein moiety indicates the pH of the sample.

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1 18. The method of claim 17, wherein the first
2 fluorescent protein moiety is a functional engineered

- 3 protein substantially identical to the amino acid sequence of Aequora green fluorescent protein (SEQ ID NO:X).
- 19. The method of claim 18, wherein the amino acid
 sequence of the protein includes the substitutions
 S65G/S72A/T203Y/H231L in the amino acid sequence of Aequora green fluorescent protein (SEQ ID NO:__).
- 1 20. The method of claim 18, wherein the amino acid 2 sequence of the protein includes the substitutions 3 S65G/V68L/Q69K/S72A/T203Y in the amino acid sequence of Aeguora green fluorescent protein (SEQ ID NO:).
- 1 21. The method of claim 18, wherein the amino acid 2 sequence of the protein includes the substitutions 3 K26R/F64L/S65T/Y66W/N146I/M153T/V163A/N164H/H231L in the 4 amino acid sequence of Aequora green fluorescent protein (SEQ ID NO:).
- 1 22. The method of claim 14, wherein the amino acid 2 sequence of the protein includes the substitution H148G in 3 the amino acid sequence of Aequora green fluorescent protein (SEO ID NO:).
- 1 23. The method of claim 14, wherein the amino acid 2 sequence of the protein includes the substitution H148Q in 3 the amino acid sequence of Aequora green fluorescent protein (SEQ ID NO:).
- 1 24. The method of claim 17, wherein the sample is a biological tissue.

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1 25. The method of claim 17, wherein the sample is a cell or a region thereof.

- 1 26. The method of claim 17, further comprising
 2 contacting the sample with a second fluorescent protein
 3 moiety whose emission intensity changes as pH varies from
 4 5 to 10, wherein the second fluorescent protein moiety emits
 5 at a second wavelength that is distinct from the first
 6 wavelength,
- 7 exciting the second protein moiety;
 8 determining the intensity of light emitted by the
 9 second protein moiety at the second wavelength; and
 10 comparing the fluorescence at the second wavelength
 to the fluorescence at the first wavelength.
- 27. The method of claim 26, wherein the second
 fluorescent protein moiety is a functional engineered
 protein substantially identical to the amino acid sequence of Aequora green fluorescent protein (SEQ ID NO:X).
- 28. The method of claim 17, wherein the first
 fluorescent protein moiety is linked to a targeting sequence.
- 29. The method of claim 28, wherein the targeting
 sequence directs the first fluorescent protein moiety to a region of a cell.
- 30. The method of claim 29, wherein the region of the cell is the cytosol, the endoplasmic reticulum, the mitochondrial matrix, the chloroplast lumen, the medial
 - trans-Golgi cisternae, the lumen of a lysosome, or the lumen of an endosome.

1 31. The method of claim 30, wherein the targeting 2 sequence comprises the amino terminal 81 amino acids of 3 human type II membrane-anchored protein qalactosyltransferase (SEQ ID NO:).

- 1 32. The method of claim 30, wherein the targeting
 2 sequence comprises the amino terminal 12 amino acids of the
 3 presequence of subunit IV of cytochrome c oxidase (SEQ ID
 4 NO:). comprises the amino terminal 81 amino acids of human
 5 type II membrane-anchored protein galactosyltransferase (SEQ
- 1 33. The method of claim 32, wherein the targeting sequence further comprises the sequence Arg-Ser-Gly-Ile.

ID NO:).

- 34. The method of claim 28, wherein the targeting
 sequence causes the first fluorescent protein moiety to be secreted from the cell.
- 35. The method of claim 17, wherein the first fluorescent protein moiety has a pKa greater than 6.1.
- 1 36. The method of claim 17, wherein the first fluorescent protein moiety has a pKa greater than 6.3.
- 1 37. The method of claim 17, wherein the first fluorescent protein moiety has a pKa greater than 6.9.
- 38. The method of claim 17, wherein the first fluorescent protein moiety has a pKa greater than 7.3.
- 1 39. The method of claim 17, wherein the first fluorescent protein moiety has a pKa greater than 7.8.

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40. The method of claim 17, wherein the first fluorescent protein moiety has a pKa of 8.0.

- 41. A method of determining the pH of a region of a 1 2 cell comprising:
- introducing into the cell a polynucleotide encoding 3 a polypeptide including an indicator having a first 4
- fluorescent protein moiety whose emission intensity changes 5
- as pH varies between 5 and 10; 6

1

- 7 culturing the cell under conditions that permit expression of the polynucleotide; 8
- 9 exciting the indicator; and
- determining the intensity of the light emitted by 10
- 11 the first protein moiety at a first wavelength, wherein the emission intensity of the first fluorescent protein moiety 12
- indicates the pH of the region of the cell in which the 13 indicator is present.
 - 42. The method of claim 41, wherein the polypeptide 1 encoded by the polynucleotide further includes a targeting 2 sequence linked by a peptide bond to the indicator.
 - 43. The method of claim 41, wherein the amino acid 1 sequence of the first fluorescent protein moiety includes 2
 - 3 the substitutions S65G/S72A/T203Y/H231L in the amino acid
 - sequence of Aequora green fluorescent protein (SEQ ID
 - NO:).
 - 44. The method of claim 41, wherein the amino acid 1 sequence of the first fluorescent protein moiety includes 2
 - 3 the substitutions S65G/V68L/O69K/S72A/T203Y in the amino
 - 4 acid sequence of Aequora green fluorescent protein (SEQ ID NO:__).

1 45. The method of claim 41, wherein the amino acid 2 sequence of the first fluorescent protein moiety includes 3 the substitutions 4 K26R/F64L/S65T/Y66W/N146I/M153T/V163A/N164H/H231L in the 5 amino acid sequence of Aequora green fluorescent protein (SEQ ID NO:__).

- 1 46. The method of claim 41, wherein the amino acid 2 sequence of the first fluorescent protein moiety includes 3 the substitution H148G in the amino acid sequence of Aequora green fluorescent protein (SEQ ID NO:__).
- 47. The method of claim 41, wherein the amino acid sequence of the first fluorescent protein moiety includes the substitution H148Q in the amino acid sequence of Aequora green fluorescent protein (SEQ ID NO:).
- 1 48. The method of claim 42, wherein the amino acid 2 of the targeting sequence comprises the amino terminal 81 amino acids of human type II membrane-anchored protein galactosyltransferase (SEQ ID NO:).
- 1 49. The method of claim 42, wherein the targeting 2 sequence comprises the amino terminal 12 amino acids of the 3 presequence of subunit IV of cytochrome c oxidase (SEQ ID NO:__).
- 50. A kit useful for the detection of the pH in a sample, the kit comprising carrier means containing one or more containers comprising a first container containing a polynucleotide comprising a nucleotide sequence encoding a functional engineered fluorescent protein whose amino acid sequence is substantially identical to the amino acid

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7 sequence of Aeguora green fluorescent protein (SEQ ID NO:__) 8 and whose emission intensity changes as pH varies between 5 and 10.

- 51. The kit of claim 50, wherein the amino acid sequence of the protein includes the substitutions S65G/S72A/T203Y/H231L in the amino acid sequence of Aequora green fluorescent protein (SEQ ID NO:).
- 52. The kit of claim 50, wherein the amino acid
 sequence of the protein includes the substitutions
 S65G/V68L/Q69K/S72A/T203Y in the amino acid sequence of
 Aequora green fluorescent protein (SEQ ID NO:).
- 1 53. The kit of claim 50, wherein the amino acid 2 sequence of the protein includes the substitutions 3 K26R/F64L/S65T/Y66W/N146I/M153T/V163A/N164H/H231L in the 4 amino acid sequence of Aequora green fluorescent protein (SEQ ID NO:).
- 1 54. The kit of claim 50, wherein the amino acid 2 sequence of the protein includes the substitution H148G in 3 the amino acid sequence of Aequora green fluorescent protein (SEQ ID NO:).
- 55. The kit of claim 50, wherein the amino acid sequence of the protein includes the substitution H148Q in the amino acid sequence of Aequora green fluorescent protein (SEC ID NO:).
- 56. A kit useful for the detection of the pH in a sample, the kit comprising carrier means containing one or more containers comprising a first container containing a

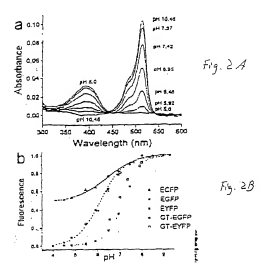
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4 functional engineered fluorescent protein whose amino acid

- 5 sequence is substantially identical to the amino acid
- 6 sequence of Aequora green fluorescent protein (SEQ ID NO:)
- $7\,$ and whose emission intensity changes as pH varies between $5\,$ and 10.

kz EGFP/EYFP cytosol/nucleus
b. GT ECFP/EGFP/EYFP Golgi: GT-ECFP/
KZ COXIV ECFP/EYFP Milochondria:

Fig. 1





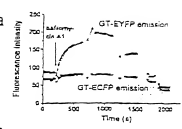
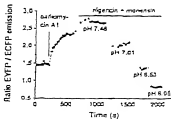


Fig. 3B



pH titration with nigericin + monensin Mitochondrial matrix pH of HeLa cells expressing YFP H148G 1000 Time (s) - H 5 μМ СССР (uncoupler) 500 6.8 8.2--8.7 -9.2 Mitochondrial matrix pH

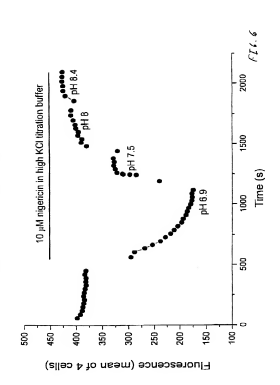
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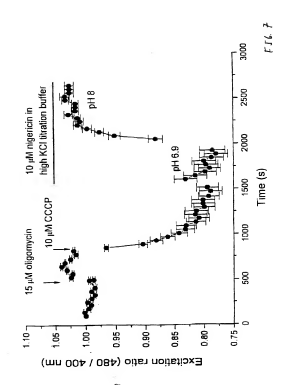
pH titration with 10 μM nigericin in high KCI 2500 2000 expressing YFP H148G in the mitochondrial matrix Milochondrial pH of chick sketetal myotubes 1500 Time (s) 66 forskolin 500 25 µM 8.0 7.0-7.6 8.9 8.2 7.8 7.4 7.2. 8.4

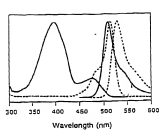
Mitochondrial matrix pH (mean of 4 cells)

Mitochondrial pH measurement in HeLa cells expressing YFP H148Q targeted to the mitochondrial matrix

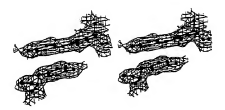


Ratiometric measurement of mitochondrial pH by expression of YFP H148G (pH sensitive) and GFP T203I (pH insensitive) in mitochondria of HeLa cells





E 16.8



F \$ 6.9